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(54) Title: DNA EXPRESSION SYSTEMS BASED ON ALPHAVIRUSES

(57) Abstract

Efficient protein production from cloned DNA in animal cells has been hampered by the lack of suitable expression systems. The requirements of such an expression system are (1) to produce functional or immunogenic forms of protein molecules in a wide variety of animal cells, (2) high efficiency and (3) technical simplicity. The present invention is related to a technical solution to this problem. A DNA molecule encoding protein sequences is inserted into engineered variants of the cDNA of a positive stranded RNA virus genome from alphavirus which then, via RNA transcription and transfection into tissue culture cells, is used to produce either chimaeric virus particles for immunization or recombinant virus for protein production. Because of optimized conditions of transfection and the nature of the virus replication the present system combines both simplicity and safety in terms of handling, efficiency in terms of level of protein and RNA production, as well as broad host range.

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DNA EXPRESSION SYSTEMS BASED ON ALPHAVIRUSES

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The present invention is related to DNA expression systems based on alphaviruses, which systems can be used to transform animal cells for use in the production of desired products, such as proteins and vaccines, in high yields.

The rapid development of biotechnology is to a large extent due to the introduction of recombinant DNA technique, which has revolutionized cellbiological and medical research by opening new approaches to elucidate the molecular mechanisms of the cell. With the aid of the techniques of cDNA cloning, large numbers of interesting protein molecules are characterized each year. Therefore, a lot of research activity is today directed to elucidate the relationship between structure and function of these molecules. Eventually this knowledge will increase our possibilities to preserve healthiness and combat diseases in both humans and animals. Indeed, there is today a growing list of new "cloned" protein products that are already used as pharmaceuticals or diagnostics.

In the recombinant DNA approaches to study biological questions, DNA expression systems are crucial elements. Thus, efficient DNA expression systems, which are simple and safe to use, give high yields of the desired product and can be used in a variety of host cells, especially also in mammalian cells, are in great demand.

Many attempts have been made to develop DNA expression systems, which fulfill these requirements. Often, viruses have been used as a source of such systems. However, up to date none of the existing viral expression systems fulfill all these requirements in a satisfying way. For instance, the <u>Baculovirus</u> expression system for cDNA is extremely efficient but can be used only in insect cells (see Reference 1 of the list of cited references; for the sake of convenience, in the following the cited references are only identified by the number they have on said list). As many important molecules will have to be produced and processed in

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cells of mammalian origin in order for them to become active, this system cannot be used in such cases. Furthermore, the Baculovirus cDNA expression system is not practically convenient for analysis of the relationship between structure and function of a protein because this involves in general the analysis of whole series of mutant variants. Today it takes about 6-8 weeks to construct a single Baculo recombinant virus for phenotype analyses. This latter problem is also true for the rather efficient Vaccinia recombinant virus and other contemporary recombinant virus cDNA expression systems (2,3). The procedure to establish stably transformed cell lines is also a very laborious procedure, and in addition, often combined with very low levels of protein expression.

Hitherto, most attempts to develop viral DNA expression systems have been based on viruses having DNA genomes or retroviruses, the replicative intermediate of the latter being double stranded DNA.

Recently, however, also viruses comprising RNA genomes have been used to develop DNA expression systems.

In EP 0 194 809 RNA transformation vectors derived from (+) strand RNA viruses are disclosed which comprise capped viral RNA that has been modified by insertion of exogenous RNA into a region non-essential for replication of said virus RNA genome. These vectors are used for expression of the function of said exogenous RNA in cells transformed therewith. The RNA can be used in solution or packaged into capsids. Furthermore, this RNA can be used to generate new cells having new functions, i.e. protein expression. The invention of said reference is generally claimed as regards host cells, (+) strand RNA viruses and the like. Nevertheless, it is obvious from the experimental support provided therein that only plant cells have been transformed and in addition only Bromo Mosaic virus, a plant

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virus, has been used as transformation vector.

Although it is stated in said reference that it is readily apparent to those skilled in the art to convert any RNA virus-cell system to a useful expression system for exogenous DNA using principals described in the reference, this has not been proven to be true in at least the case of animal cell RNA viruses. The reasons for this seem to be several. These include:

 Inefficiencies in transfecting animal cells with in vitro transcribed RNA;

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- 2) Inefficiency of apparently replication competent RNA transcripts to start RNA replication after commonly used transfection procedures;
- 3) The inability to produce high titre stocks of recombinant virus that does not contain any helper virus;
- 4) The inability to establish stable traits of transformed cells expressing the function of the exogenous RNA.

In Proc. Natl. Acad. Sci. USA, Vol 84, 1987, pp 4811-4815 a gene expression system based on a member of the Alphavirus genus, viz. Sindbis virus, is disclosed which is used to express the bacterial CAT (chloramphenical acetyltransferase) gene in avian cells, such as chicken embryo fibroblasts.

Xiong et al., Science, Vol 243, 1989, 1188-1191 also disclose a gene expression system based on Sindbis virus. This system is said to be efficient in a broad range of animal cells. Expression of the bacterial CAT gene in insect, avian and mammalian cells inclusive of human cells is disclosed therein.

Even though it is known from prior art that one

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member of the Alphavirus genus, the Sindbis virus, can tolerate insertion and direct the expression of at least one foreign gene, the bacterial chloramfenicol acetyl transferase (CAT) gene, it is evident from the results described that both systems described above are both ineffective in terms of exogenous gene expression and also very cumbersome to use. Hence, neither system has found any usage in the field of DNA expression in animal cells today.

In the first example a cDNA copy of a defective interfering (DI) virus variant of Sindbis virus was used to carry the CAT gene. RNA was transcribed in vitro and used to transfect avian cells and some CAT protein production could be demonstrated after infecting cells with wild-type Sindbis virus. The latter virus provided the viral replicase for expression of the CAT construct. The inefficiency of this system depends on 1) low level of initial DI-CAT RNA transfection (0.05-0.5 % of cells) and 2) inefficient usage of the DI-CAT RNA for protein translation because of unnatural and suboptimal protein intitation translation signals. This same system also results in packaging of some of the recombinant DI-CAT genomes into virus particles. However, this occurs simultaneously with a very large excess of wild-type Sindbis virus production. Therefore, the usage of this mixed virus stock for CAT expression will be much hampered by the fact that most of the replication and translation activity of the cells infected with such a stock will deal with the wild-type and not with recombinant gene expression.

Much of the same problems are inherent to the other Sindbis expression system described. In this an RNA replication competent Sindbis DNA vector is used to carry the CAT gene. RNA produced in vitro is shown to replicate in animal cells and CAT activity is found. However, as only a very low number of cells are transfected the overall CAT production remains low. Another

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possible explanation for this is that the Sindbis construct used is not optimal for replication. Wild-type Sindbis virus can be used to rescue the recombinant genome into particles together with an excess of wildtype genomes and this mixed stock can then be used to express a CAT protein via infection. However, this stock has the same problems as described above for the recombinant DI system. The latter paper shows also that if virus is amplified by several passages increased titres of the recombinant virus particles can be obtained. However, one should remember that the titre of the wild-type virus will increase correspondingly and the original problem of mostly wild-type virus production remains. There are also several potential problems when using several passages to produce a mixed virus stock. As there is no selected pressure for preservation of the recombinant genomes these might easily 1) undergo rearrangements and 2) become outnumbered by wild-type genomes as a consequence of less efficient replication and/or packaging properties.

Another important aspect of viral DNA expression vectors is use thereof to express antigens of unrelated pathogens and thus they can be used as vaccines against such pathogens.

Development of safe and effective vaccines against viral diseases has proven to be quite a difficult task. Although many existing vaccines have helped to combat the worldwide spread of many infectious diseases, there is still a large number of infectious agents against which effective vaccines are missing. The current procedures of preparing vaccines present several problems:

(1) it is often difficult to prepare sufficiently large amounts of antigenic material; (2) In many cases there is the additional hazard that the vaccine preparation is not killed or sufficiently attenuated; (3) Effective vaccines are often hard to produce since there is a major difficulty in presenting the antigenic epitope in

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an immunologically active form; (4) In the case of many viruses, genetic variations in the antigenic components results in the evolution of new strains with new serological specificities, which again creates a need for the development of new vaccines.

Two types of viral DNA vectors have been developed in order to overcome many of these problems in vaccine production. These either provide recombinant viruses or provide chimaeric viruses. The recombinant viruses contain a wild-type virus package around a recombinant genome. These particles can be used to infect cells which then produce the antigenic protein from the recombinant genome. The chimaeric viruses also contain a recombinant genome but this specifies the production of an antigen, usually as part of a normal virus structural protein, which then will be packaged in progeny particles and e.g. exposed on the surface of the viral spike proteins. The major advantages of these kind of virus preparations for the purpose of being used as a vaccine are 1) that they can be produced in large scale and 2) that they provide antigen in a natural form to the immunological system of the organism. Cells, which have been infected with recombinant viruses, will synthesize the exogenous antigen product, process it into peptides that then present them to T cells in the normal way. In the case of the chimaeric virus there is, in addition, an exposition of the antigen in the context of the subunits of the virus particle itself. Therefore, the chimaeric virus is also called an epitope carrier.

The major difficulty with these kind of vaccine preparations are, how to ensure a safe and limited replication of the particles in the host without side effects. So far, some success has been obtained with vaccinia virus as an example of the recombinant virus approach (69) and of polio virus as an example of a chimaeric particle (70-72). As both virus variants are

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based on commonly used vaccine strains one might argue that they could be useful vaccine candidates also as recombinant respectively chimaeric particles (69-72). However, both virus vaccines are combined with the risk for side effects, even severe ones, and in addition these virus strains have already been used as vaccines in large parts of the population in many countries.

As is clear from the afore mentioned discussion there is much need to develop improved DNA expression systems both for an easy production of important proteins or polypeptides in high yields in various kinds of animal cells and for the production of recombinant viruses or chimaeric viruses to be used as safe and efficient vaccines against various pathogenes.

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Thus, an object of the present invention is to provide an improved DNA expression system based on virus vectors which can be used both to produce proteins and polypeptides and as recombinant virus or chimaeric virus, which system offers many advantages over prior art.

To that end, according to the present invention there is provided an RNA molecule derived from an alphavirus RNA genome and capable of efficient infection of animal host cells, which RNA molecule comprises the complete alphavirus RNA genome regions, which are essential to replication of the said alphavirus RNA, and further comprises an exogenous RNA sequence capable of expressing its function in said host cell, said exogenous RNA sequence being inserted into a region of the RNA molecule which is non-essential to replication thereof.

Alphavirus is a genus belonging to the family Togaviridae having single stranded RNA genomes of positive polarity enclosed in a nucleocapsid surrounded by an evelope containing viral spike proteins.

The Alphavirus genus comprises among others the Sindbis virus, the Semliki Forest virus (SFV) and the Ross River virus, which are all closely related.

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According to a preferred embodiment of the invention, the Semliki Forest virus (SFV) is used as the basis of the DNA expression system.

The exogenous RNA sequence encodes a desired genetic trait, which is to be conferred on the virus or the host cell, and said sequence is usually complementary to a DNA or cDNA sequence encoding said genetic trait. Said DNA sequence may be comprised of an isolated natural gene, such as a bacterial or mammalian gene, or may constitute a synthetic DNA sequence coding for the desired genetic trait i.e. expression of a desired product, such as an enzyme, hormone, etc. or expression of a peptide sequence defining an exogenous antigenic epitope or determinant.

If the exogenous RNA sequence codes for a product, such as a protein or polypeptide, it is inserted into the viral RNA genome replacing deleted structural protein encoding region(s) thereof, whereas a viral epitope encoding RNA sequence may be inserted into structural protein encoding regions of the viral RNA genome, which essentially do not comprise deletions or only have a few nucleosides deleted.

The RNA molecule can be used per se, e.g. in solution to transform animal cells by conventional transfection, e.g. the DEAE-Dextran method or the calcium phosphate precipitation method. However, the rate of transformation of cells, and, thus the expression rate can be expected to increase substantially if the cells are transformed by infection with infectious viral particles. Thus, a suitable embodiment of the invention is related to an RNA virus expression vector comprising the RNA molecule of this invention packaged into infectious particles comprising the said RNA within the alphavirus nucleocapsid and surrounded by the membrane including the alphavirus spike proteins.

The RNA molecule of the present invention can be packaged into such particles without restraints pro-

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vided that it has a total size corresponding to the wild type alphavirus RNA genome or deviating therefrom to an extent compatible with package of the said RNA into the said infectious particles.

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These infectious particles, which include recombinant genomes packaged to produce a pure, high titre recombinant virus stock, provides a means for exogenous genes or DNA sequences to be expressed by normal virus particle infection, which as regards transformation degree, is much more efficient than RNA transfection.

According to a suitable embodiment of the invention such infectious particles are produced by cotransfection of animal host cells with the present RNA which lacks part of or the complete region(s) encoding the structural viral proteins together with a helper RNA molecule transcribed in vitro from a helper DNA vector comprising the SP6 promoter region, those 5' and 3' regions of the alphavirus cDNA which encode cis acting signals needed for RNA replication and the region encoding the viral structural proteins but lacking essentially all of the nonstructural virus proteins encoding regions including sequenses encoding RNA signals for packaging of RNA into nucleocapsid particles, and culturing the host cells.

According to another aspect of the invention efficient introduction of the present RNA into animal host cells can be achieved by electroporation. For example, in the case of Baby Hamster Kidney (BHK) cells a transformation degree of almost 100 % has been obtained for the introduction of an RNA transcript derived from SFV cDNA of the present invention. This makes it possible to reach so high levels of exogenous protein production in every cell that the proteins can be followed in total cell lysates without the need of prior concentration by antibody precipitation.

By electroporation, it is also possible to obtain a high degree of cotransfection in the above process for WO 92/10578 PCT/SE91/00855

production of infectious particles comprising packaged RNA of the present invention. Essentially all animal cells will contain both the present RNA molecule and the helper RNA molecule, which leads to a very efficient trans complementation and formation of infectious partcles. A pure recombinant virus stock, consisting of up to 109-1010 infectious particles, can be obtained from 5 x 106 cotransfected cells after only a 24 h incubation. Furthermore, the so obtained virus stock is very safe to use, since it is comprised of viruses containing only the desired recombinant genome, which can infect host cells but can not produce new progeny virus.

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Theoretically, a regeneration of a wild-type virus genome could take place when producing the recombinant virus in the contransfected cells. However, the possibility to avoid spread of such virus can be eliminated by incorporating a conditionally lethal mutation into the structural part of the helper genome. Such a mutation is described in the experimental part of this application. Thus, the virus produced with such a helper will be noninfectious if not treated in vitro under special conditions.

The technique of electroporation is well known within the field of biotechnology and optimal conditions can be established by the man skilled in the art. For instance, a BioRad Gene pulser apparatus (BioRad, Richmond, CA, USA) can be used to perform said process.

The RNA molecule of the present invention is derived by in vivo or in vitro transcription of a cDNA clone, originally produced from an alphavirus RNA and comprising an inserted exogenous DNA fragment encoding a desired genetic trait.

Accordingly, the present invention is also related to a DNA expression vector comprising a full-length or partial cDNA complementary to alphavirus RNA or parts thereof and located immediately downstream of the SP6

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RNA polymerase promoter and having a 5'ATGG, a 5'GATGG or any other 5' terminus and a TTTCCA₆₉ACTAGT or any other 3' terminus.

According to one aspect of the present invention portions of the viral cDNA are deleted, the deletions comprising the complete or part of the region(s) encoding the virus structural proteins, and the vector further comprises an integrated polylinker region, which may correspond to BamHI-SmaI-XmaI, inserted at a location which enables an exogenous DNA fragment encoding a foreign polypeptide or protein to be inserted into the vector cDNA for subsequent expression in an animal host cell.

According to another aspect of this invention, the vector is comprised of full-length cDNA wherein an exogenous DNA fragment encoding a foreign epitopic peptide sequence can be inserted into a region coding for the viral structural proteins.

It is appreciated that this cDNA clone with its exogenous DNA insert is very efficiently replicated after having been introduced into animal cells by transfection.

A very important aspect of the present invention is that it is applicable to a broad range of host cells of animal origin. These host cells can be selected from avian, mammalian, reptilian, amphibian, insect and fish cells. Illustrative of mammalian cells are human, monkey, hamster, mouse and porcine cells. Suitable avian cells are chicken cells, and as reptilian cells viper cells can be used. Cells from frogs and from mosquitoes and flies (Drosophila) are illustrative of amphibian and insecticidal cells, respectively. A very efficient virus vector/host cell system according to the invention is based on SFV/BHK cells, which will be discussed more in detail further below.

However, even though a very important advantage of the present DNA expression vector is that it is very

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efficient in a broad variety of animal cells it can also be used in other eucaryotic cells and in procaryotic cells.

The present invention is also related to a method to produce transformed animal host cells comprising transfection of the cells with the present RNA molecule or with the present transcription vector comprised of cDNA and carrying an exogenous DNA fragment. According to a suitable embodiment of the invention, transfection is produced by the above mentioned electroporation method, a very high transfection rate being obtained.

A further suitable transformation process is based on infection of the animal host cells with the above mentioned infectious viral particles comprising the present RNA molecule.

The transformed cells of the present invention can be used for different purposes.

One important aspect of the invention is related to use of the present transformed cells to produce a polypeptide or a protein by culturing the transformed cells to express the exogenous RNA and subsequent isolation and purification of the product formed by said exepression. The transformed cells can be produced by infection with the present viral particles comprising exogenous RNA encoding the polypeptide or protein as mentioned above, or by transfection with an RNA transcript obtained by in vitro transcription of the present DNA vector comprised of cDNA and carrying an exogenous DNA fragment coding for the polypeptide or the protein.

Another important aspect of the invention is related to use of the present transformed cells for the production of antigens comprised of chimaeric virus particles for use as immunizing component in vaccines or for immunization purposes for in vivo production of immunizing components for antisera production.

Accordingly, the present invention is also related to an antigen consisting of a chimaeric alphavirus having

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an exogenous epitopic peptide sequence inserted into its structural proteins.

Preferably, the chimaeric alphavirus is derived from SFV.

According to a suitable embodiment, the exogenous epitopic peptide sequence is comprised of an epitopic peptide sequence derived from a structural protein of a virus belonging to the immunodeficiency virus class inclusive of the human immunodeficiency virus types.

A further aspect of the invention is related to a vaccine preparation comprising the said antigen as immunizing component.

In said vaccine the chimaeric alphavirus is suitably attenuated by comprising mutations, such as the conditionally lethal SFV-mutation described before, amber (stop codon) or temperature sensitive mutations, in its genome.

For instance, if the chimaeric virus particles containing the afore mentioned conditional lethal mutation in its s tructural proteins (a defect to undergo a certain proteolytical cleavage in host cell during morphogenesis) is used as a vaccine then this is first activated by limited proteolytic treatment before given to the organism so that it can infect recipient cells. New chimaeric particles will be formed in cells infected with the activated virus but these will again be of the lethal phenotype and further spread of infection is not possible.

The invention is also concerned with a method for the production of the present antigen comprising

- a) in vitro transcription of the cDNA of the present DNA vector carrying an exogenous DNA fragment encoding the foreign epitopic peptide sequence and transfection of animal host cells with the produced RNA transcript, or
- b) transfection of animal host cells with the said cDNA of the above step a),

culturing the transfected cells and recovering the chimaeric alphavirus antigen. Preferably, transfection is produced by electroporation.

Still another aspect of the invention is to use a recombinant virus containing exogenous RNA encoding a polypeptide antigen for vaccination purpose or to produce antisera. In this case the recombinant virus or the conditionally lethal variant of it is used to infect cells in vivo and antigen production will take place in the infectious cells and used for antigen presentation to the immunological system.

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According to another embodiment of the invention, the present antigen is produced in an organism by using in vivo infection with the present infectious particles containing exogenous RNA encoding an exogenous epitopic peptide sequence.

In the following, the present invention will be illustrated more in detail with reference to the Semliki Forest virus (SFV), which is representative for the alphaviruses. This description can be more fully understood in conjunction with the accompanying drawings in which:

Fig. 1 is a schematic view over the main assembly and disassembly events involved in the life cycle of the Semliki Forest virus, and also shows regulation of the activation of SFV entry functions by p62 cleavage and pH;

Fig. 2 illustrates the use of translocation signals during synthesis of the structural proteins of SFV; top, the gene map of the 26S subgenomic RNA; middle, the process of membrane translocation of the p62, 6K and E1 proteins; small arrows on the lumenal side denote signal peptidase cleavages; at the bottom, the characteristics of the three signal peptides are listed;

Fig. 3 shows features that make SFV an excellent

choice as an expression vector;

Fig. 4 A-C show the construction of full-length infectious clones of SFV; Fig. 4A shows a schematic restriction map of the SFV genome; primers used for initiating cDNA synthesis are indicated as arrows, and the cDNA inserts used to assemble the final clone are showed as bars; Fig. 4B shows plasmid pPLH211, i.e. the SP6 expression vector used as carrier for the full-length infectious clone of SFV, and the resulting plasmid pSP6-SFV4; Fig. 4C shows the structure of the SP6 promoter area of the SFV clone; the stippled bars indicate the SP6 promoter sequence, and the first necleotide to be transcribed is marked by an asterisk; underlined regions denote authentic SFV sequences;

Fig. 5 shows the complete nucleotide sequence of the pSP6-SFV4 RNA transcript as DNA (U=T) and underneath the DNA sequence, the amino acid sequence of the non-structural polyprotein and the structural polyprotein;

Fig. 6 shows an SFV cDNA expression system for the production of virus after transfection of in vitro made RNA into cells;

Fig. 7 shows the construction of the SFV expression vectors pSFV1-3 and of the Helper 1;

Fig. 8 shows the polylinker region of SFV vector plasmids pSFV1-3; the position of the promoter for the subgenomic 26S RNA is boxed, and the first nucleotide to be transcribed is indicated by an asterisk;

Fig. 9 is a schematic presentation of in vivo packaging of pSFV1-dhfr RNA into infectious particles using helper trans complementation; (dhfr means di-hydrofolate reductase)

Fig. 10 shows the use of trypsin to convert p62-containing noninfectious virus particles to infectious particles by cleavage of p62 to E2 and E3;

Fig. 11 shows the expression of heterologous proteins in BHK cells upon RNA transfection by electroporation; and

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Fig. 12 shows in its upper part sequences encompassing the major antigenic site of SFV and the in vitro made substitutions leading to a BamHI restriction endonuclease site, sequences spanning the principal neutralizing domain of the HIV gp120 protein, and the HIV domain inserted into the SFV carrier protein E2 as a BamHI oligonucleotide; and its lower part is a schematic presentation of the SFV spike structure with blow-ups of domain 246-251 in either wild type or chimaeric form.

The alphavirus Semliki Forest virus (abbreviated SFV in the following text) has for some 20 years been used as model system in both virology and cell biology to study membrane biosynthesis, membrane structure and membrane function as well as protein-RNA interactions (4, 5). The major reason for the use of SFV as such a model is due to its simple structure and efficient replication.

With reference to Fig. 1-3, in the following the SFV and its replication are explained more in detail. In essential parts, this disclosure is true also for the other alphaviruses, such as the Sindbis virus, and many of the references cited in this connection are indeed directed to the Sindbis virus. SFV consists of an RNAcontaining nucleocapsid and a surrounding membrane composed of a lipid bilayer and proteins, a regularly arranged icosahedral shell of a protein called C protein forming the capsid inside which the genomic RNA is packaged. The capsid is surrounded by the lipid bilayer that contains three proteins called E1, E2, and E3. These so-called envelope proteins are glycoproteins and their glycosylated portions are on the outside of the lipid bilayer, complexes of these proteins forming the "spikes" that can be seen in electron micrographs to project outward from the surface of the virus.

The SFV genome is a single-stranded 5'-capped and 3'-polyadenylated RNA molecule of 11422 nucleotides (6,7).

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It has positive polarity, i.e. it functions as an mRNA, and naked RNA is able to start an infection when introduced into the cytoplasm of a cell. Infection is initiated when the virus binds to protein receptors on the host cell plasma membrane, whereby the virions become selectively incorporated into "coated pits" on the surface of the plasma membrane, which invaginate to form coated vesicles inside the cell, whereafter said vesicles bearing endocytosed virions rapidly fuse with organelles called endosomes. From the endosome, the virus escapes into the cell cytosol as the bare nucleocapsid, the viral envelope remaining in the endosome. Thereafter, the nucleocapsid is "uncoated" and, thus, the genomic RNA is released. Referring now to Fig. 1, infection then proceeds with the translation of the 5' two-thirds of the genome into a polyprotein which by self-cleavage is processed to the four nonstructural proteins nsP1-4 (8). Protein nsP1 encodes a methyl transferase which is responsible for virus-specific capping activity as well as initiation of minus strand synthesis (9, 10); nsP2 is the protease that cleaves the polyprotein into its four subcomponents (11, 12); nsP3 is a phosphoprotein (13, 14) of as yet unknown function, and nsP4 contains the SFV RNA polymerase activity (15, 16). Once the nsP proteins have been synthesized they are responsible for the replication of the plus strand (42S) genome into full-length minus strands. These molecules then serve as templates for the production of new 42S genomic RNAs. They also serve as templates for the synthesis of subgenomic (265) RNA. This 4073 nucleotides long RNA is colinear with the last one-third of the genome, and its synthesis is internally initiated at the 26S promoter on the 42S minus strands (17, 18).

The capsid and envelope proteins are synthesized in different compartments, and they follow separate pathways through the cytoplasm, viz. the envelope proteins

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are synthesized by membrane-bound ribosomes attached to the rough endoplasmic reticulum, and the capsid protein is synthesized by free ribosomes in the cytosol. However, the 26S RNA codes for all the structural proteins of the virus, and these are synthesized as a polyprotein precursor in the order C-E3-E2-6K-E1 (19). Once the capsid (C) protein has been synthesized it folds to act as a protease cleaving itself off the nascent chain (20, 21). The synthesized C proteins bind to the recently replicated genomic RNA to form new nucleocapsid structures in the cell cytoplasm.

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The said cleavage reveals an N-terminal signal sequence in the nascent chain which is recognized by the signal recognition particle targeting the nascent chain - ribosome complex to the endoplasmic reticulum (ER) membrane (22, 23), where it is cotranslationally translocated and cleaved by signal peptidase to the three structural membrane proteins p62 (precursor form of E3/E2), 6K and E1 (24, 25). The translocational signals used during the synthesis of the structural proteins are illustrated in Fig. 2. The membrane proteins undergo extensive posttranslational modifications within the biosynthetic transport pathway of the cell. The p62 protein forms a heterodimer with E1 via its E3 domain in the endoplasmic reticulum (26). This dimer is transported out to the plasma membrane, where virus budding occurs through spike nucleocapsid interactions. At a very late (post-Golgi) stage of transport the p62 protein is cleaved to E3 and E2 (27), the forms that are found in mature virions. This cleavage activates the host cell binding function of the virion as well as the membrane fusion potential of E1. The latter activity is expressed by a second, low-pH activation step after the virus enters the endosomes of a new host cell and is responsible for the release of the viral nucleocapsid into the cell cytoplasm (28-32). The mature virus particles contain one single copy of the RNA

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genome encapsidated within 180 copies of the capsid protein in a T=3 symmetry, and is surrounded by a lipid bilayer carrying 240 copies of the spike trimer protein consisting of E1+E2+E3 arranged in groups of three in a T=4 symmetry (33).

The SFV entry functions are activated and regulated by p62 cleavage and pH. More specifically, the p62-E1 heterodimers formed in the ER are acid resistant. When these heterodimers are transported to the plasma membrane via the Golgi complex the E1 fusogen cannot be activated in spite of the mildly acidic environment, since activation requires dissociation of the complex. As is illustrated in Fig. 1, the released virus particles contain E2E1 complexes. Since the association between E2 and E1 is sensitive to acidic pH, during entry of the virus into a host cell through endocytosis the acidic milieu of the endosome triggers the dissociation of the spike complex (E1 E2 E3) resulting in free E1. The latter can be activated for the catalysis of the fusion process between the viral and endosomal membranes in the infection process as disclosed above.

As indicated in the preceding parts of the disclosure, the alphavirus system, and especially the SFV system, has several unique features which are to advantage in DNA expression systems. These are summarized below with reference to Fig. 3.

- 1. Genome of positive polarity. The SFV RNA genome is of positive polarity, i.e. it functions directly as mRNA, and infectious RNA molecules can thus be obtained by transcription from a full-length cDNA copy of the genome.
- 2. Efficient replication. The infecting RNA molecule codes for its own RNA replicase, which in turn drives an efficient RNA replication. Indeed, SFV is one of the most efficiently replicating viruses known. Within a few hours up to 200.000 copies of the plus-RNAs are made in a single cell. Because of the abundance of

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these molecules practically all ribosomes of the infected cell will be enrolled in the synthesis of the virus encoded proteins, thus overtaking host protein synthesis (34), and pulse-labelling of infected cells results in almost exclusive labelling of viral proteins. During a normal infection 10⁵ new virus particles are produced from one single cell, which calculates to at least 10⁸ protein molecules encoded by the viral genome (5).

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3. Cytoplasmic replication. SFV replication occurs in the cell cytoplasm, where the virus replicase transcribes and caps the subgenomes for production of the structural proteins (19). It would obviously be very valuable to include this feature in a cDNA expression system to eliminate the many problems that are encountered in the conventional "nuclear" DNA expression systems, such as mRNA splicing, limitations in transcription factors, problems with capping efficiency and mRNA transport.

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4. Late onset of cytopathic effects. The cytopathic effects in the infected cells appear rather late during infection. Thus, there is an extensive time window from about 4 hours after infection to up to 24 hours after infection during which a very high expression level of the structural proteins is combined with negligible morphological change.

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5. Broad host range. This phenomenon is probably a consequence of the normal life cycle which includes transmission through arthropod vectors to wild rodents and birds in nature. Under laboratory conditions, SFV infects cultured mammalian, avian, reptilian and insect cells (35) (Xiong, et al, loc. cit.)

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6. In nature SFV is of very low pathogenicity for humans. In addition, the stock virus produced in tissue culture cells is apparently apathogenic. By means of specific mutations it is possible to create conditionally lethal mutations of SFV, a feature that is of

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great use to uphold safety when massproduction of virus stocks is necessary.

In the nucleotide and amino acid sequences the following abbreviations have been used in this specification:

Ala, alanine; lle, isoleucine; leu, leucine; Met, methionine; Phe, phenylalanine; Pro, proline; Trp, tryptophan; Val, valine; Asn, asparagine; Cys, cysteine; Gln, glutamine; Gly, glycine; Ser, serine; Thr, threonine; Tys, tyrosine; Arg, arginine; His, histidine; Lys, lysine; Asp, aspartic acid; Glu, glutamic acid; A, adenine; C, cytosine; G, guanine; T, thymine; U, uracil.

The materials and the general methodology used in the following examples are disclosed below.

- Materials. Most restriction enzymes, DNA Polymerase I, Klenow fragment, calf intestinal phosphatase, T4 DNA ligase and T4 Polynucleotide kinase were from Boehringer (Mannheim, FRG). SphI, StuI and KpnI together with RNase inhibitor (RNasin) and SP6 Polymerase were from Promega Biotec (Madison, WI). Sequenase (Modified T7 polymerase) was from United States Biochemical (Cleveland, Ohio). Proteinase K was from Merck (Darmstadt, FRG). Ribonucleotides, deoxyribonucleotides, dideoxyribonucleotides and the cap analogue $m^7G(5')ppp(5')G$ were from Pharmacia (Sweden). Oligonucleotides were produced using an Applied Biosystems synthesizer 380B followed by HPLC and NAP-5 (Pharmacia) purification. Spermidine, phenylmethylsulfonyl fluoride (PMSF), diethylpyrocarbonate (DEPC), bovine serum albumin (BSA), creatine phosphate and creatine phosphokinase were from Sigma (St. Louis, Mo). Pansorbin was from CalBiochem (La Jolla, CA). Agarose was purchased from FMC BioProducts (Rockland, Maine), and acrylamide from BioRad (Richmond, CA). $L-[^{35}S]$ methionine and $\alpha-[^{35}S]$ -dATP- α -S were from Amersham.
 - 2. Virus growth and purification: BHK-21 cells were

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grown in BHK medium (Gibco Life Technologies, Inc., New York) supplemented with 5 % fetal calf serum, 10 % tryptose phosphate broth, 10 mM HEPES (N-2-hydroxy-ethylpiperazine-N'-2-ethanesulfonic acid) and 2 mM glutamine. 90 % confluent monolayers were washed once with PBS and infected with SFV in MEM containing 0.2 % bovine serum albumin (BSA), 10 mM HEPES and 2 mM glutamine at a multiplicity of 0.1. Twenty-four hours post infection (p.i.) the medium was collected and cell debris removed by centrifugation at 8,000 xg for 20 min at 4°C. The virus was pelleted from the medium by centrifugation at 26,000 rpm for 1.5 h in an SW28 rotor at 4°C. The virus was resuspended in TN containing 0.5 mM EDTA.

3. Metabolic labeling and immunoprecipitation. Confluent monolayers of BHK cells grown in MEM supplemented with 10 mM HEPES, 2 mM glutamine, 0.2 % BSA, 100 IU/mol of penicillin and 100 μ g/ml streptomycin, were infected at a multiplicity of 50 at 37°C. After 1 h p.i. the medium was replaced with fresh and growth continued for 3.5 h. The medium was removed and cells washed once with PBS and overlayed with methionine-free MEM containing 10 mM HEPES and 2 mM glutamine. After 30 min at 37°C the medium was replaced with the same containing 100 μ Ci/ml of [35S]methionine (Amersham) and the plates incubated for 10 min at 37°C. The cells were washed twice with labeling medium containing 10X excess methionine and then incubated in same medium for various times. The plates were put on ice, cells washed once with ice-cold PBS and finally lysis buffer (1 % NP-40 - 50 mM Tris-HCl, pH 7.6 - 150 mM NaCl - 2 mM EDTA) containing 10 μ g/ml PMSF (phenylmethylsulfonyl fluoride) was added. Cells were scraped off the plates. and nuclei removed by centrifugation at 6,000 rpm for 5 min at 4°C in an Eppendorf centrifuge. Immunoprecipitations of proteins was performed as described (31). Briefly, antibody was added to lysate and the mixture

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kept on ice for 30 min. Complexes were recovered by binding to Pansorbin for 30 min on ice. Complexes were washed once with low salt buffer, once with high salt buffer, and once with 10 mM Tris-HCl, pH 7.5, before heating with gel loading buffer. To precipitate dhfr, SDS was added to 0.1 % and the mixture heated to 95°C for 2 min followed by addition of 10 volumes of lysis buffer. Anti-E1 [8.139], anti-E2 [5.1] (36), and anti-C [12/2] (37) monoclonals have been described. The human transferrin receptor was precipitated with the monoclonal antibody OKT-9 in ascites fluid. This preparation was provided by Thomas Ebel at our laboratory using a corresponding hybridoma cell line obtained from ATCC (American Typ Culture Collection) No CRL 8021. Polyclonal rabbit anti-mouse dhfr was a kind gift from E. Hurt (European Molecular Biology Laboratory, Heidelberg, FRG) and rabbit anti-lysozyme has been described (38).

4. Immunofluorescence. To perform indirect immunofluorescence, infected cell monolayers on glass coverslips were rinsed twice with phosphate-buffered saline (PBS) and fixed in -20°C methanol for 6 min. After fixation, the methanol was removed and the coverslip washed 3 times with PBS. Unspecific antibody binding was blocked by incubation at room temperature with PBS containing 0.5 % gelatin and 0.25 % BSA. The blocking buffer was removed and replaced with same buffer containing primary antibody. After 30 min at room temperature the reaction was stopped by washing 3 times with PBS. Binding of secondary antibody (FITC-conjugated sheep anti-mouse [BioSys, Compiégne, France]) was done as for the primary antibody. After 3 washes with PBS and one rinse with water the coverslip was allowed to dry before mounting in Moviol 4-88 (Hoechst, Frankfurt am Main, FRG) containing 2.5 % DABCO (1,4-diazobicyclo-[2.2.2]-octane).

5. DNA procedures. Plasmids were grown in Escherichia

coli DH5α (Bethesda Research Laboratories) [recA endAl gyrA96 thil hsdR17 supE44 relAl Δ(lacZYA-argF)U169 φ80dlacZΔ(M15)]. All basic DNA procedures were done essentially as described (39). DNA fragments were isolated from agarose gels by the freeze-thaw method (40) including 3 volumes of phenol during the freezing step to increase yield and purity. Fragments were purified by benzoyl-naphthoyl-DEAE (BND) cellulose (Serva Feinbiochemica, Heidelberg, FRG) chromatography (41). Plasmids used for production of infectious RNA were purified by sedimentation through 1 M NaCl followed by banding in CsCl (39). In some cases plasmids were purified by Qiagen chromatography (Diagen Gmbh, Düsseldorf, FRG).

6. Site-directed oligonucleotide mutagenesis. For

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oligonucleotide mutagenesis, relevant fragments of the SFV cDNA clone were subcloned into M13mp18 or mp 19 (42) and transformed (43) into DH5αFIQ [endA1 hsdR1 supE44 thi1 recA1 gyrA96 reIA1 φ80dlacΔ(M15) Δ(lacZYAargF)U169/F'proAB laclq lacZA(M15) Tn 5] (Bethesda Research Laboratories). RF DNA from these constructs was transformed into RZ1032 (44) [Hfr KL16 dut1 ung1 thi1 relA1 supE44 zbd279:Tn10.], and virus grown in the presence of uridine to incorporate uracil residues into the viral genome. Single stranded DNA was isolated by phenol extraction from PEG precipitated phage. Oligonucleotides were synthesized on an Applied Biosystems 380B synthesizer and purified by gel filtration over NAP-5 columns (Pharmacia). The oligonucleotides 5'-CGGCCAGTGAATTCTGATTGGATCCCGGGTAATTAATTGAATTACATCCC-TACGCAAACG, 5'-GCGCACTATTATAGCACCGGCTCCCGGGTAATTAATT-GACGCAAACGTTTTACGGCCGCCGG and 5'-GCGCACTATTATAGCACCATG-GATCCGGGTAATTAATTGACGTTTTACGGCCGCCGGTGGCG were used to insert the new linker sites [BamHI-SmaI-XmaI] into the SFV cDNA clone. The oligonucleotides 5'-CGGCGGTCCTA-GATTGGTGCG and 5'-CGCGGGCGCCACCGGCGGCCG were used as sequencing primers (SP1 and SP2) up- and downstream of

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the polylinker site. Phosphorylated oligonucleotides were used in mutagenesis with Sequenase (Unites States Biochemicals, Cleveland, Ohio) as described earlier (44, 45). In vitro made RF forms were transformed into DH5 α F'IQ and the resulting phage isolates analyzed for the presence of correct mutations by dideoxy sequencing according to the USB protocol for using Sequenase. Finally, mutant fragments were reinserted into the full-length SFV cDNA clone. Again, the presence of the appropriate mutations was verified by sequencing from the plasmid DNA. Deletion of the 6K region has been described elsewhere.

7. In vitro transcription. SpeI linearized plasmid DNA was used as template for in vitro transcription. RNA was synthesized at 37°C for 1 h in 10-50 μ l reactions containing 40 mM Tris-HCl (pH 7.6), 6 mM spermidine-HCl, 5 mM dithiothreitol (DTT), 100 μ g/ml of nuclease free BSA, 1 mM each of ATP, CTP and UTP, 500 μ M of GTP, 1 unit/ μ l of RNasin and 100-500 units/ml of SP6 RNA polymerase. For production of capped transcripts (46), the analogs m7G(5')ppp(5')G or m⁷G(5')ppp(5')A were included in the reaction at 1 mM. For quantitation of RNA production, trace amounts of $[\alpha^{-32}P]$ -UTP (Amersham) was included in the reactions and incorporation measured from trichloroacetic acid precipitates. When required, DNA or RNA was digested at 37°C for 10 min by adding DNase 1 or RNase A at 10 units/ μg template or 20 μ g/ml respectively.

8. RNA transfection. Transfection of BHK monolayer cells by the DEAE-Dextran method was done as described previously (47). For transfection by electroporation, RNA was added either directly from the in vitro transcription reaction or diluted with transcription buffer containing 5 mM DTT and 1 unit/ μ l of RNasin. Cells were trypsinized, washed once with complete BHK-cell medium and once with ice-cold PBS (without MgCl₂ and CaCl₂) and finally resuspended in PBS to give 10⁷ cells/ml. Cells

were either used directly or stored (in BHK medium) on ice over night. For electroporation, 0.5 ml of cells were transferred to a 0.2 cm cuvette (BioRad), 10-50 μ l of RNA solution added and the solution mixed by inverting the cuvette. Electroporation was performed at room temperature by two consecutive pulses at 1.5 kV/25 μF using a BioRad Gene Pulser apparatus with its pulse controller unit set at maximum resistance. After incubation for 10 min, the cells were diluted 1:20 in complete BHK-cell medium and transferred onto tissue culture plates. For plaque assays, the electroporated cells were plated together with about 3x105 fresh cells per ml and incubated at 37°C for 2 h, then overlayed with 1.8 % low melting point agarose in complete BHKcell medium. After incubation at 37°C for 48 h, plagues were visualized by staining with neutral red.

9. Gel electrophoresis. Samples for sodium dodecyl sulfate - polyacrylamide gel electrophoresis (SDS-PAGE) were prepared and run on 12 % separating gels with a 5 % stacking gel as previously described (48). For resolving the 6K peptide, a 10 % - 20 % linear acrylamide gradient gel was used. Gels were fixed in 10 % acetic acid - 30 % methanol for 30 min before exposing to Kodak XAR-5 film. When a gel was prepared for fluorography (49), it was washed after fixation for 30 min in 30 % methanol and then soaked in 1M sodium salicylate - 30 % methanol for 30 min before drying. Nucleic acids were run on agarose gels using 50 mM Tris-borate - 2.5 mM Na₂EDTA as buffer. For staining 0.2 µg/ml of ethidium bromide was included in the buffer and gel during the run.

Example 1

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In this example a full-length SFV cDNA clone is prepared and placed in a plasmid containing the SP6 RNA polymerase promoter to allow in vitro trancription of full-length and infectious transcripts. This plasmid which is designated pSP6-SFV4 has been deposited on 28

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European Collection of Animal Cell Cultures, Porton Down, Salisbury, Wiltshire, U.K:, and given the provisional accession number 91112826.

As illustrated in Fig. 4A-C the strategy for construction the SFV clone was to prime cDNA synthesis on several positions along the template RNA downstream of suitable restriction endonuclease sites defined by the known nucleotide sequence of the SFV RNA molecule. Virus RNA was isolated by phenol-chloroform extraction from purified virus (obtainable among others from the Arbovirus collection in Yale University, New Haven, USA) and used as template for cDNA synthesis as previously described (50). First strand synthesis was primed at three positions, using 5'-TTTCTCGTAGTTCTCCTC-GTC as primer-1 (SFV coordinate 2042-2062) and 5'-GTTA-TCCCAGTGGTTGTTCTCGTAATA as primer-2 (SFV coordinate 3323-3349) and an oligo-dT₁₂₋₁₈ as primer -3 (3' end of SFV) Fig. 4A).

Second strand synthesis was preceded by hybridization of the oligonucleotide 5'-ATGGCGGATGTGTGACATACACGACGCC (identical to the 28 first bases of the genome sequence of SFV) to the first strand cDNA. After completion of second strand synthesis cDNA was trimmed and in all cases except in the case of the primer-1 reaction, the double-stranded adaptor 5'-AATTCAAGCTTGCGGCCGCACTAGT / GTTCGAACGCCGGCGTGATCA-3' (5'-sticky-EcoRI-HindIII-NotI-XmaIII-SpeI-blunt-3') was added and the cDNa cloned into EcoRl cleaved pTZ18R (Pharmacia, Sweden) as described (51). The cloning of the 5' end region was done in a different way. Since SFV contains a HindIII site at position 1947, cDNA primed with primer-1 should contain this area and therefore HindIII could be used to define the 3' end of that cDNA. To obtain a restriction site at the very 5' end of the SFV, cDNA was cloned into Smal-HindIII cut pGEM1 (Promega Biotec.,

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Madison, W1). Since the SFV genome starts with the sequence 5'-ATGG, ligation of this onto the blunt CCC-3' end of the Smal site created an Ncol site C'CATGG. Although the SFV sequence contains 3 NcoI sites, none of these are within the region preceding the HindIII site, and thus these 5' end clones could be further subcloned as NcoI-HindIII fragments into a vector especially designed for this purpose (see below). The original cDNA clones in pGEM1 were screened by restriction analysis and all containing inserts bigger than 1500 bp were selected for further characterization by sequencing directly from the plasmid into both ends of the insert, using SP6 or T7 sequencing primers. The SFV 5'-end clones in pTZ18R were sequenced using lac sequencing primers. To drive in vitro synthesis of SFV RNA the SP6 promoter was used. Cloning of the SFV 5' end in front of this promoter without adding too many foreign nucleotides required that a derivative of pGEM1 had to be constructed. Hence, pGEM1 was opened at EcoRl and Bal31 deletions were created, the DNA blunted with T4 DNA polymerase and an Ncol oligonucleotide (5'-GCCATGGC) added. The clones obtained were screened by colony hybridization (39) with the oligonucleotide 5'-GGTGACACTATAGCCATGGC designed to pick up (at suitable stringency) the variants that had the NcoI sequence immediately at the transcription initiation site of the SP6 promoter (G underlined). Since the Bal31 deletion had removed all restriction sites of the multicloning site of the original plasmid, these were restored by cloning a PvuI-NcoI fragment from the new variant into another variant of pGEM1 (pDH101) that had an NcoI site inserted at its HindIII position in the polylinker. This created the plasmid pDH201. Finally, the adaptor used for cloning the SFV cDNA was inserted into pDH201 between the EcoRI and PvuII sites to create plasmid pPLH211 (Fig. 4B). This plasmid was then used as recipient for SFV cDNA fragments in the assembly of the

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full-length clone by combining independent overlapping subclones using these sites. The fragments and the relevant restriction sites used to assemble the fulllength clone, pSP6-SFV4, are depicted in (Fig. 4A). For the 5'-end, the selected fragment contained the proper SFV sequence 5'-ATGG, with one additional G-residue in front. When this G-residue was removed it reduced transcription efficiency from SP6 but did not affect infectivity of the in vitro made RNA. Thus, the clone used for all subsequent work contains the G-residue at the 5' end. For the 3'-end of the clone, a cDNA fragment containing 69 A-residues was selected. By inclusion of the unique SpeI site at the 3'-end of the cDNa, the plasmid can be linearized to allow for runoff transcription in vitro giving RNA-carrying 70 Aresidues. Fig. 4C shows the 5' and 3' border sequences of the SFV cDNA clone. The general outline how to obtain and demonstrate infectivity of the full-length SFV RNA is depicted in Fig. 6. The complete nucleotide sequence of the pSP6-SFV4 SP6 transcript together with the amino acid sequences of the nonstructural and the structural polyproteins is shown in Fig. 5.

Typically, about 5 µg of RNA per 100 ng of template was obtained using 10 units of polymerase, but the yield could be increased considerably by the use of more enzyme. The conditions slightly differ from those reported earlier for the production of infectious transcripts of alphaviruses (52) (47). A maximum production of RNA was obtained with rNTP concentrations at 1 mM. However, since infectivity also is dependent on the presence of a 5'cap structure optimal infectivity was obtained when the GTP concentration in the transcription reaction was halved. This drop had only a marginal effect on the amounts of RNA produced but raised the specific infectivity by a factor of 3 (data not shown).

The cDNA sequence shown in Fig. 5 has been used in the following examples. However, sequences having one

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or a few nucleotides, which differ from those shown in Fig. 5, could also be useful as vectors, even if these might be less efficient as illustrated above with the SFV cDNA sequence lacking the first 5'-G nucleotide in Fig. 5.

Example 2.

In this example the construction of SFV DNA expression vectors is disclosed.

The cDNA clone coding for the complete genome of SFV obtained in Example 1 was used to construct a SFV DNA expression vector by deletion of the coding region of the 26S structural genes to make way for heterologous inserts. However, the nonstructural coding region, which is required for the production of the nsP1-4 replicase complex is preserved. RNA replication is dependent on short 5' (nt 1-247) (53, 54, 55) and 3' (nt 11423-11441) sequence elements (56, 57), and therefore, also these had to be included in the vector construct, as had the 26S promoter just upstream of the C gene (17, 18).

As is shown in Fig. 7, first, the XbaI (6640)-NsiI (8927) fragment from the SFV cDNA clone pSP6-SFV4 from Example 1 was cloned into pGEM7Zf(+) (Promega Corp., Wl, USA) (Step A). From the resulting plasmid, pGEM7Zf(+)-SFV, the EcoRI fragment (SFV coordinates 7391 and 88746) was cloned into M13mp19 to insert a BamHI - XmaI Smal polylinker sequence immediately downstream from the 26S promoter site using site-directed mutagenesis (step B). Once the correct mutants had been verfied by sequencing from M13 ssDNA (single stranded), the EcoRI fragments were reinserted into pGEM7Zf(+)-SFV (step C) and then cloned back as XbaI-Nsl fragments into pSP6-SFV4 (step D). To delete the major part of the cDNA region coding for the structural proteins of SFV, these plasmids were then cut with AsuII (7783) and NdeI (11033), blunted using Klenow fragment in the presence of all four nucleotides, and religated to create the

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final vectors designated pSFV1, pSFV2 and pSFV3, respectively (step E). The vectors retain the promoter region of the 26S subgenomic RNA and the last 49 amino acids of the E1 protein as well as the complete non-coding 3' end of the SFV genome.

In the vectors the subgenomic (26S) protein coding portion has been replaced with a polylinker sequence allowing the insertional cloning of foreign cDNA sequences under the 26S promoter. As is shown in Fig. 8 these three vectors have the same basic cassette inserted downstream from the 26S promoter, i.e. a polylinker (BamHI-SmaI-XmaI) followed by a translational stop-codons in all three reading frames. The vectors differ as to the position where the polylinker cassette has been inserted. In pSFV1 the cassette is situated 31 bases downstream of the 26S transcription initiation site. The initiation motive of the capsid gene translation is identical to the consensus sequence (58). Therefore, this motive has been provided for in pSFV2, where it is placed immediately after the motive of the capsid gene. Finally, pSFV3 has the cassette placed immediately after the initiation codon (AUG) of the capsid gene. Sequencing primers (SP) needed for checking both ends of an insert have been designed to hybridize either to the 26S promoter region (SP1), or to the region following the stop codon cassette (SP2).

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Note that the 26S promoter overlaps with the 3'-end of the nsP4 coding region. For psFV2, the cloning site is positioned immediately after the translation initiation site of the SFV capsid gene. For psFV3, the cloning site is positioned three nucleotides further downstream, i.e. immediately following to the initial AUG codon of the SFV capsid gene. The three translation stop codons following the polylinker are boxed. The downstream sequencing primer (SP1) overlaps with the 26S promoter, and the upstream sequencing primer (Sp2)

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overlaps the XmaIII site.

Example 3

In this example an in vivo packaging system encompassing helper virus vector constructs is prepared. .

The system allows SFV variants defective in structural protein functions, or recombinant RNAs derived from the expression vector construct obtained in Example 2, to be packaged into infectious virus particles. Thus, this system allows recombinant RNAs to be introduced into cells by normal infection. The helper vector, called pSFV-Helper1, is constructed by . deleting the region between the restriction endonuclease sites AccI (308) and AccI (6399) of pSP6-SFV4 obtained in Example 1 by cutting and religation as shown in Fig. 7, step F. The vector retains the 5' and 3' signals needed for RNA replication. Since almost the complete nsP region of the Helper vector is deleted, RNA produced from this construct will not replicate in the cell due to the lack of a functional replicase complex. As is shown in Fig. 9, after transcription in vitro of pSFV1-recombinant and helper cDNAs, helper RNA is cotransfected with the pSFV1 - recombinant derivative, the helper construct providing the structural proteins needed to assemble new virus particles, and the recombinant providing the nonstructural proteins needed for RNA replication, SFV particles comprising recombinant genomes being produced. The cotransfection is preferably produced by electroporation as is disclosed in Example 6 and preferably BHK cells are used as host cells.

To package the RNA a region at the end of nsP1 is required, an area which has been shown to bind capsid protein (57, 59). Since the Helper lacks this region, RNA derived from this vector will not be packaged and hence, transfections with recombinant and Helper produces only virus particles that carry recombinantderived RNA. It follows that these viruses cannot be

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passaged further and thus provide a one-step virus stock. The advantage is that infections with these particles will not produce any viral proteins.

Example 4

This example illustrates the construction of variants of the full-length SFV cDNA clone from Example 1 that allow insertion of foreign DNA sequences encoding foreign epitopes, and the production of recombinant (chimaeric) virus carrying said foreign epitopes as integral parts of the p62, E2 or E1 spike proteins.

To this end, a thorough knowledge of the function, topology and antigenic structure of the E2 and E1 envelope proteins has been of the essence. Earlier studies on the pathogenicity of alphaviruses have shown that antibodies against E2 are type-specific and have good neutralizing activity while those against E1 generally are group-specific and are nonneutralizing (5). However, not until recently have antigenic sites of the closely related alphaviruses SFV, Sindbis, and Ross River been mapped and correlated to the level of amino acid sequence (60, 61, 62, 63). These studies have shown that the most dominant sites in question are at amino acid positions 216, 234 and 246-251 of the SFV E2 spike protein. Interestingly, these three sites are exactly the same as the ones predicted by computer analysis. In the present example domain 246-251 was used, since this area has a highly conserved structure and hydropathy profile within the group of alphaviruses. Insertion of a gene encoding a foreign epitope into the 246-251 region of the pSP6-SFV4 p62 protein yields particles with one new epitope on each heterodimer, i.e. 240 copies.

To create a unique restriction endonuclease site that would allow specific insertion of foreign epitopes into the E2 portion of the SFV genome, a BamHI site was inserted by site directed mutagenesis using the oligonucleotide 5'-GATCGGCCTAGGAGCCCGAGAGCCC.

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Example 5

In this example a conditionally lethal variant of SFV is constructed from the SFV cDNA obtained in Example 1, which variant carries a mutation in the p62 protein resulting in a noncleavable from of said protein, with the result that this variant as such cannot infect new host cells, unless first cleaved with exogenously added protease.

As illustrated in Fig. 10, this construct can be advantageously used as a vaccine carrier for foreign epitopes, since this form of the virus cannot enter new host cells although assembled with wild type efficiency in transfected cells. The block can be overcome by trypsin treatment of inactive virus particles. This converts the particle into a fully entry-competent form which can be used for amplification of this virus variant stock.

Once activated the SFV variant will enter cells normally through the endocytic pathway and start infection. Viral proteins will be made and budding takes place at the plasma membrane. However, all virus particles produced will be of inactive form and the infection will thus cease after one round of replication. The reason for the block in infection proficiency is a mutation which has been introduced by site directed mutagenesis into the cleavage site of p62. This arginine to leucine substitution (at amino acid postion 66 of the E3 portion of the p62 protein) changes the consensus features of the cleavage site so that it will not be recognized by the host cell proteinase that normally cleaves the p62 protein to the E2 and E3 polypeptides during transport to the cell surface. Instead, only exogenously added trypsin will be able to perform this cleavage, which in this case occurs at the arginine residue 65 immediately preceding the original cleavage site. As this cleavage regulates the activation of the entry function potential of the

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virus by controlling the binding of the entry spike subunit, the virus particle carrying only uncleaved p62 will be completely unable to enter new host cells.

The creation of the cleavage deficient mutation E2 has been described earlier (29). An Asull - Ns \(\lambda \) fragment spanning this region was then isolated and cloned into the full-length cDNA clonepSP6-SFV4.

Example 6

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In this example transfection of BHK cells with SFV RNA molecules transcribed in vitro from full-length cDNA from Example 1 or variants thereof or the SFV vectors from Example 2, which comprise exogenous DNA, is disclosed. The transfection is carried out by electroporation which is shown to be very efficient at optimized conditions.

BHK cells were transfected with the above SFV RNA molecules by electroporation and optimal conditions were determined by varying parameters like temperature, voltage, capacitance, and number of pulses. Optimal transfection was obtained by 2 consecutive pulses of 1.5 kV at 25 μ F, under which negligible amounts of cells were killed. It was found that it was better to keep the cells at room tempeature than at 0°C during the whole procedure. Transfection by electroporation was also measured as a function of input RNA. As expected, an increase in transfection frequency was not linearly dependent on RNA concentration, and about 2 μ g of cRNA were needed to obtain 100 % transfection.

On comparison with conventional transfection, this is a great improvement. For example, with DEAE-Dextran transfection optimally, only 0.2 % of the cells were transfected:

Example 7

This example illustrates heterologous gene expression driven by the SFV vector, pSFV1 from Example 2, for genes encoding the 21 kD cytoplasmic mouse dihydrofolate reductase (dhfr), the 90 kD membrane protein

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human transferrin receptor (TR), and finally the 14 kD secretory protein chicken lysozyme. The dhfr gene was isolated from pGEM2-dhfr (64) as a BamHI-HindIII fragment blunted with Klenow fragment and inserted into SmaI-cut pSFV1. The transferrin receptor gene was first cloned from pGEM1-TR (64, 65) as an XbaI-EcoRI fragment into pGEM7ZF(+) and subsequently from there as a BamHI fragment into pSFV1. Finally, a BamHI fragment from pGEM2 carrying the lysozyme gene (21) was cloned into pSFV1.

To study the expression of the heterologous proteins, in vitro-made RNA of the dhfr and TR constructs was electroporated into BHK cells. RNA of wild type SFV was used as control. At different time points post electroporation (p.e.) cells were pulse-labeled for 10 min followed by a 10 min chase, whereafter the lysates were analyzed by gel electrophoresis and autoradiography. The results are shown in Figure 11. More specifically, BHK cells were transfected with RNAs of wild type SFV, pSFV1-dhfr, and pSFV1-TR, pulse-labeled at 3, 6, 9, 12, 15 and 24 h p.e. Equal amounts of lysate were run on a 12 % gel. The 9 h sample was also used in immunoprecipitation (IP) of the SFV, the dhfr and the transferrin receptor proteins. Cells transfected with pSFV1lysozyme were pulse-labeled at 9 h p.e. and then chased for the times (hours) indicated. An equal portion of lysate or medium was loaded on the 13,5 % gel. IP represents immunoprecipitation from the 1 h chase lysate sample. The U-lane is lysate of labeled but untransfected cells. At 3 h p.e.hardly any exogenous proteins were made, since the incoming RNA starts with minus strand synthesis which does not peak until about 4-5 h p.e. (5). At this time point, almost all labeled proteins were of hos origin. In contrast, at 6 h p.e. the exogenous proteins were synthesized with great efficiency, and severe inhibition of host protein synthesis was evident. This was even more striking at 9 h

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p.e., when maximum shut down had been reached. Efficient production of the heterologous proteins continued up to 24 h p.e., after which production slowed down (data not shown), indicating that the cells had entered a stationary phase.

Since chicken lysozyme is a secretory protein, its expression was analyzed both from cell lysates and from the growth medium. Cells were pulse-labeled at 9 h p.e. and then chased up to 8 h. The results are shown in Fig. 11. Although lysozyme was slowly secreted, almost all labeled material was secreted to the medium during the chase.

Example 8

This example illustrates the present in vivo packaging system.

In vitro-made RNA of pSFV1-TR was mixed with Helper RNA at different ratios and these mixtures were cotransfected into BHK cells. Cells were grown for 24 h after which the culture medium was collected and the virus particles pelleted by ultracentrifugation. The number of infectious units (i.u.) was determined by immunofluorescence. It was found that a 1:1 ratio of Helper and recombinant most efficiently produced infectious particles, and on the average 5 x 106 cells yielded 2.5×10^9 i.u. The infectivity of the virus stock was tested by infecting BHK cells at different multiplicaties of infection (m.o.i.). In Fig. 11 the results for expression of human transferrin receptor in BHK cells after infection by such in vivo packaged particles carrying pSFV1-TR recombinant RNA is shown to the lower right. 200 μ l of virus diluted in MEM (including 0,5'% BAS and 2 mM glutamine) was overlaid on cells to give m.o.i. values ranging from 5 to 0.005. After 1 h at 37°C, complete BHK medium was added and growth continued for 9 h, at which time a 10 min pulse (100 μ Ci ³⁵S-methionine/ml) and 10 min chase was performed, and the cells dissolved in lysis buffer. 10

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 μ l out of the 300 μ l lysate (corresponding to 30,000 cells) was run on the 10 % gel, and the dried gel was exposed for 2 h at -70°C. Due to the high expression level, only 3,000 cells are needed to obtain a distinct band on the autoradiograph with an over night exposure.

Thus, it was found that efficient protein production and concomitant hos protein shut-off occurred at about 1 i.u. per cell. Since one SFV infected cell produces on the average 10⁸ capsid protein molecules, it follows that a virus stock produced from a single electroporation can be used to produce 10¹⁷ protein molecules equaling about 50 mg of protein.

From the foregoing experimental results it is obvious that the present invention is related to very useful and efficient expression system which lacks several of the disadvantages of the hitherto existing expression system. The major advantages of the present system are shortly summarized as follows:

- (1) High titre recombinant virus stocks can be produced in one day by one transfection experiment. There is no need for selection/screening, plaque purification and amplification steps. This is appreciated since an easy production of recombinant virus is especially important in experiments where the phenotypes of large series of mutants have to be characterized.
- (2) The recombinant virus stock is free from helper virus since only the recombinant genome but not the helper genome contains a packaging signal.
- (3) The recombinant virus can be used to infect the recombinant genome in a "natural" and nonleakey way into a large variety of cells including insect and most higher euoaryotic cell types. Such a wide host range is very useful for an expressions system

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especially when cell-type-specific posttranslational modification reactions are required for the activity of the expressed protein.

extremely high, the level corresponding to those of the viral proteins during infection. There is also a host cell protein shut-off which makes it possible to follow the foreign proteins clearly in cell lysates without the need for antibody mediated antigen concentration. This will facilitate DNA expression experiments in cell biology considerably. Furthermore, problems of interference by the endogenous counter part to an expressed protein (i.e. homo-oligomerization reactions) can be avoided.

Example 9

This example illustrates epitope carriers.

A very important example where vaccine development is of the utmost importance concerns the acquired immunodeficiency syndrome (AIDS) caused by the human immunodeficiency virus HIV-1 (66, 67). Sofar, all attempts to produce an efficient vaccine against HIV-1 have failed, although there was a very recent report that vaccination with disrupted SIV-1 (Simian immunodeficiency virus) to a certain extent may give protection against infections of that virus (68). However, development of safe and effective vaccine against HIV-1 will be very difficult due to the biological properties of the virus. In the present exampel one epitope of HIV-1 was inserted into an antigenic domain of the E2 protein of SFV. The epitope used is located in glycoprotein gp120 of HIV-1, spanning amino acids 309-325. This forms the variable loop of HIV-1 and is situated immediately after an N-glycosylated site.

A chimaera was constructed where the 309-325 epitope of HIV was inserted into the BamHI site using cassette

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insertion of ready-made oligonucleotides encoding the HIV epitope. The required base substitutions at the BamHI site did not lead to any amino acid changes in the vector, although two amino acids (Asp and Glu) changed places. This change did not have any deleterious effect since in vitro made vector RNA induced cell infection with wild type efficiency. Fig. 12 shows the sequences in the area of interest in the epitope carrier. In preliminary experiments, it has been shown that chimaeric proteins were produced. The proteins can be immunoprecipitated with anti-HIV anti-bodies. It is to be expected that these are also used for production of chimaeric virus particles that can be used for vaccine preparation against HIV. Such particles are shown in Fig. 12, lower part.

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Claims

- 1. An RNA molecule derived from an alphavirus RNA genome and capable of efficient infection of animal host cells, which RNA molecule comprises the complete alphavirus RNA genome regions, which are essential to replication of the said alphavirus RNA, and further comprises an exogenous RNA sequence capable of expressing its function in said host cell, said exogenous RNA sequence being inserted into a region of the RNA molecule which is non-essential to replication thereof.
- 2. The RNA of claim 1, wherein the said alphavirus is Semliki Forest virus (SFV).
- 3. The RNA of claim 1 or 2, wherein the exogenous RNA sequence encodes a protein, a polypeptide or a peptide sequence defining an exogenous antigenic epitope or determinant.
- 4. The RNA of claim 3 wherein the exogenous RNA sequence encodes an epitope sequence of a structural protein of an immunodeficiency virus inclusive of the human immunodeficiency virus (HIV) types.
- 5. The RNA of any preceding claim, wherein the alphavirus derived RNA molecule regions comprise a 5' terminal portion, the coding region(s) for non structural proteins required for RNA replication, the subgenome promoter region and a 3' terminal portion of said viral RNA.
- 6. The RNA of claim 2, 3 or 5, wherein the exogenous RNA sequence encodes a foreign polypeptide or protein and is integrated into the SFV subgenomic 26S RNA substituting deleted parts thereof.
- 7. The RNA of claim 2, 3, 4 or 5, wherein the exogenous RNA sequence encodes a foreign viral epitopic peptide sequence and is located in a region of the RNA coding for structural alphavirus proteins enabling the exogenous RNA to be expressed as said viral epitope as part of the matured virus particle.
- 8. The RNA of claim 2, 3, 4 or 5, wherein the exogenous RNA sequence encodes a foreign viral epitopic peptide sequence inserted into the p62 spike precursor subunit encoding region of the SFV genome.
- 9. An RNA expression vector comprising the RNA of any preceding claim packaged into infectious particles comprising the RNA within the alphavirus nucleocapsid and surrounded by membrane with alphavirus spike proteins.

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- 10. The vector of claim 9, wherein the RNA has a total size corresponding to the wild type alphavirus RNA genome or deviating therefrom to an extent compatible with package of the RNA into the infectious particles.
- 11. DNA transcription vector comprising a cDNA having one strand complementary to the RNA of any of claims 1 to 8.
- 12. A DNA expression vector comprising a full-length or partial cDNA complementary to alphavirus RNA or parts thereof and located immediately downstream of the SP6 RNA polymerase promoter and having a 5'ATGG or 5'GATGG or any other 5' terminus and a TTTCCA69ACTAGT or any other 3' terminus.
- 13. The vector of claim 12 having portions of the viral cDNA deleted, the deletions comprising the complete or part of the region(s) encoding the virus structural proteins, and further comprising an integrated polylinker region, which may correspond to BamHI-SmaI-XmaI, inserted at a location which enables an exogenous DNA fragment encoding a foreign polypeptide or protein to be inserted into the vector cDNA for subsequent expression in an animal host cell.
- 14. The vector of claim 12 or 13 wherein the alphavirus is SFV.
- 15. The vector of claim 12 or 14 comprising full-length cDNA and further comprising an exogenous DNA fragment encoding a foreign epitopic peptide sequence or antigenic determinant inserted into a region of the viral structural proteins.
- 16. The vector of claim 15 wherein the exogenous DNA fragment is inserted into the p62 spike precursor subunit encoding region of the SFV cDNA.
- 17. The vector of any preceding claim comprising an SFV derived cDNA which carries a conditionally lethal SFV mutation in the region encoding the p62 cleavage site, a cellularly uncleavable but extracellularly cleavable form of p62 being expressed.
- 18. The vector of claim 13 comprising SFV-derived cDNA, the vector being pSFV1, pSFV2 or pSFV3 having a structure as shown in Fig. 8.
- 19. An RNA transcript derived from transcription of the DNA-vector of any of claims 12-18 carrying an exogenous DNA fragment.
 - 20. A method to produce the vector of claim 9 or 10

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wherein the alphavirus derived RNA lacks part of or the complete region(s) encoding the structural viral proteins, the method comprising cotransfection of animal host cells with the RNA transcript of claim 19, wherein the alphavirus RNA lacks part(s) of or the complete region(s) encoding the viral structural proteins, with helper RNA transcribed in vitro from a helper DNA vector and culturing the host cells.

- 21. The method of claim 20 wherein the cotransfection is produced by electroporation of the host cells.
- 22. Helper vector for use in the method according to claim 20 or 21, said vector being comprised of the DNA vector of claim 12 wherein the regions encoding non structural virus proteins are almost completely deleted, including sequences encoding RNA signals for packaging of RNA into nucleocapsid particles, but the 5' and 3' signals needed for RNA replication and the region encoding the promoter for the structural subgenome are in addition to those encoding the structural region preserved.
- 23. Helper vector of claim 22 wherein the cDNA has its origin from SFV and the deletion extends from the AccI (308) to the AccI (6399) restriction endonuclease site of the full-length cDNA vector of claim 12.
- 24. Helper vector of claim 22 and 23 where the structural region contains the mutation described in claim 17 or another conditionally lethal mutation.
- 25. The method of claim 20 wherein cells transformed to produce helper RNA according to claims 20, 22 or 23 are transfected with RNA transcript of claim 19.
- 26. A host cell of animal origin transformed with the RNA of any of claims 1-8, the DNA transcription vector of claims 11 or the DNA vector of any of claims 12-18 carrying an exogenous DNA fragment.
- 27. The host cell of claim 26 wherein the cell is an avian, a mammalian, a reptilian, an amphibian, an insecticidal or a fish cell.
- 28. The host cell of claim 27 which is the hamster BHK cell.
- 29. A method to produce the transformed host cell of claim 26, 27 or 28 comprising transfection of the cell with the RNA of any of claim 1-8, with the cDNA of claim 11 or of any of

claims 12-18 carrying an exogenous DNA fragment or infection of the cell with the infectious viral particles of claim 9 or 10.

30. The method of claim 29 wherein the transfection is produced by electroporation of the host cell.

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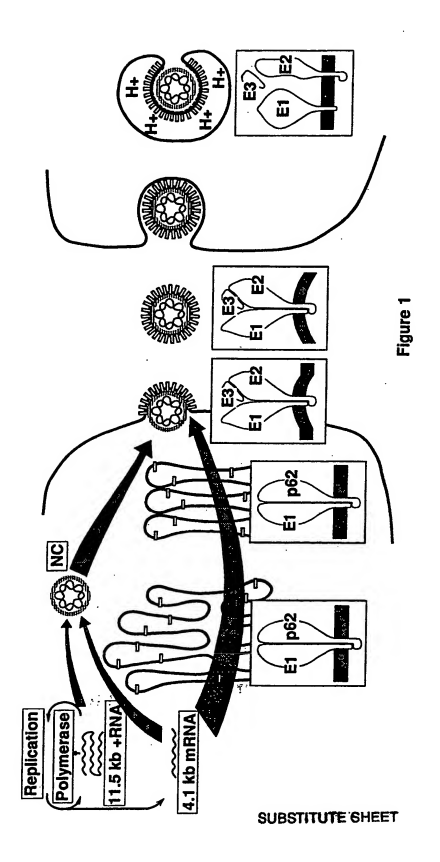
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- 31. A method for the production of a polypeptide or protein comprising infection of animal host cells with infectious particles according to claim 9 or 10, containing exogenous RNA encoding said polypeptide or protein and produced according to method of claim 20 or 21, culturing the said transformed cells to express the exogenous RNA and isolation and purification of the product formed by said expression.
- 32. A method for the production of a polypeptide or protein comprising in vitro transcription of the cDNA of the vector of any of claims 11-18 carrying an exogenous DNA fragment coding for the polypeptide or protein, transfection of animal host cells with the produced RNA transcript, transformed animal host cells being obtained harbouring the RNA transcript, culturing the said transformed cells to express the exogenous RNA and isolation and purification of the product formed by said expression.
- 33. The metod of claim 32 wherein the vector cDNA is comprised of the cDNA of the vector of claim 17 carrying the exogenous DNA fragment.
- 34. An antigen consisting of a chimaeric alphavirus having an exogenous epitopic peptide sequence or antigenic determinant inserted into its structural proteins.
- 35. The antigen of claim 34 wherein the chimaeric alphavirus is derived from SFV.
- 36. The antigen of claim 34 or 35, wherein the exogenous epitopic peptide sequence is comprised of an epitopic peptide sequence derived from a structural protein of a virus belonging to the immunodeficiency virus class inclusive of the human immunodeficiency virus types.
- 37. Vaccine preparation comprising the antigen of claim 34, 35 or 36 as immunizing component.
- 38. Vaccine of claim 37 wherein the chimaeric alphavirus is attenuated by comprising the conditionally lethal SFV mutation of claim 17, an amber (stop codon) a temperature sensitive mutation or other mutation in its genome.
 - 39. A method for the production of an antigen of claim

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- 34, 35 or 36 comprising
- a) in vitro transcription of the cDNA of the vector of any of claims 11-18 carrying an exogenous DNA fragment encoding the foreign epitopic peptide sequence or antigenic determinant and transfection of animal host cells with the produced RNA transcript, or
- b) transfection of animal host cells with the said cDNA of the above step a),
- culturing the transfected cells and recovering the chimaeric alphavirus antigen.
- 40. The method of claim 32, 33 or 39 wherein the transfection is produced by electroporation of the host cell.
- 41. A method for the production of an antigen in an organism by using in vivo infection with infectious particles according to claim 9 or 10 containing exogenous RNA encoding an exogenous epitopic peptide sequence or antigenic determinant, and produced according the claim 20 or 21.



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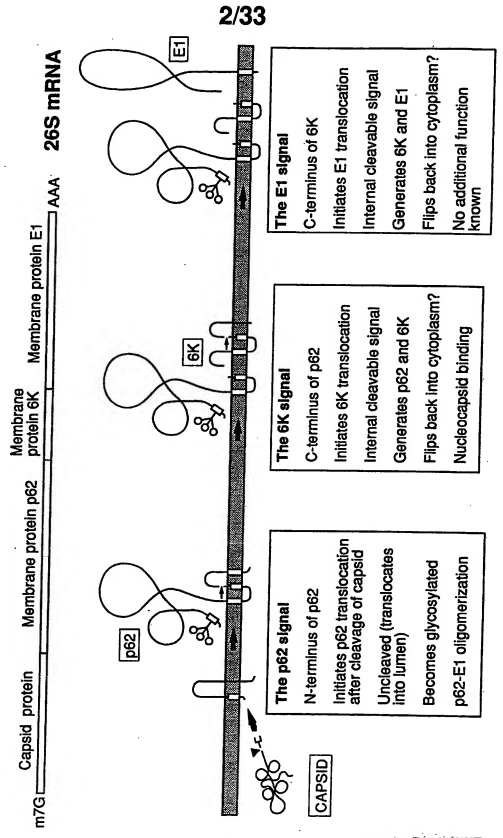
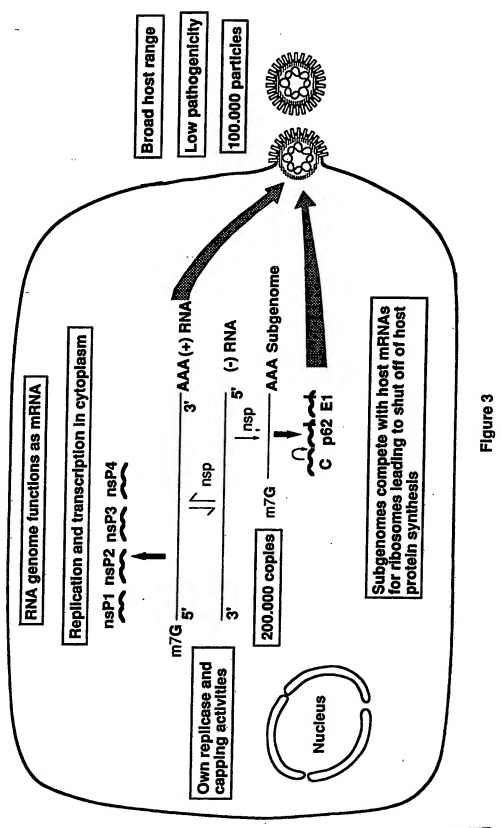


Figure 2



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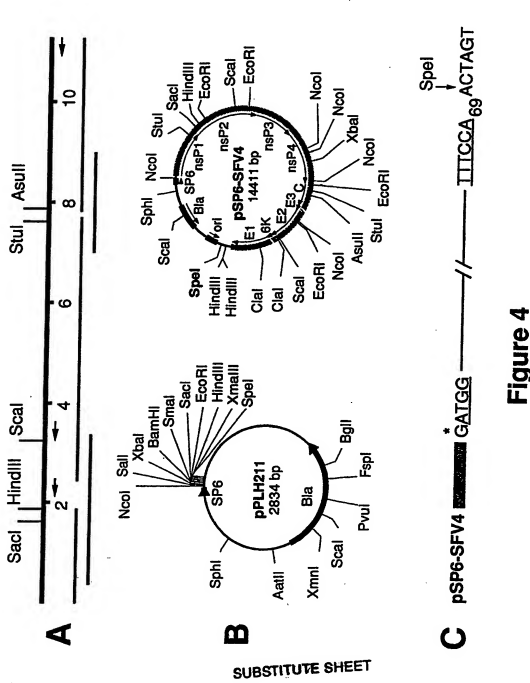


Figure 5 (1)

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GNIGGERAL GIGIGACAIA CACGACGCA AMAGAIIIIG IICCAGCICC IGCCACCICC GU												
GCTACGCGAG AGATTAACCA CCCACG ATG GCC GCC AAA GTG CAT GTT GAT ATT 113 Met Ala Ala Lys Val His Val Asp Ile 5												
GAG GCT GAC AGC CCA TTC ATC AAG TCT TTG CAG AAG GCA TTT CCG 158												
Glu Ala Asp Ser Pro Phe Ile Lys Ser Leu Gln Lys Ala Phe Pro 10 15 20												
TCG TTC GAG GTG GAG TCA TTG CAG GTC ACA CCA AAT GAC CAT GCA 203 Ser Phe Glu Val Glu Ser Leu Gln Val Thr Pro Asn Asp His Ala 25 30 35												
AAT GCC AGA GCA TTT TCG CAC CTG GCT ACC AAA TTG ATC GAG CAG 248 Asn Ala Arg Ala Phe Ser His Leu Ala Thr Lys Leu Ile Glu Gln 40 45 50												
GAG ACT GAC AAA GAC ACA CTC ATC TTG GAT ATC GGC AGT GCG CCT 293 Glu Thr Asp Lys Asp Thr Leu Ile Leu Asp Ile Gly Ser Ala Pro 55 60 65												
TCC AGG AGA ATG ATG TCT ACG CAC AAA TAC CAC TGC GTA TGC CCT 338 Ser Arg Arg Met Met Ser Thr His Lys Tyr His Cys Val Cys Pro 70 75 80												
ATG CGC AGC GCA GAA GAC CCC GAA AGG CTC GAT AGC TAC GCA AAG 383 Met Arg Ser Ala Glu Asp Pro Glu Arg Leu Asp Ser Tyr Ala Lys 85 90 95												
AAA CTG GCA GCG GCC TCC GGG AAG GTG CTG GAT AGA GAG ATC GCA 428 Lys Leu Ala Ala Ala Ser Gly Lys Val Leu Asp Arg Glu Ile Ala 100 105 110												
GGA AAA ATC ACC GAC CTG CAG ACC GTC ATG GCT ACG CCA GAC GCT 473 Gly Lys Ile Thr Asp Leu Gln Thr Val Met Ala Thr Pro Asp Ala 115 120 125												
GAA TCT CCT ACC TTT TGC CTG CAT ACA GAC GTC ACG TGT CGT ACG 518 Glu Ser Pro Thr Phe Cys Leu His Thr Asp Val Thr Cys Arg Thr 130 135 140												
GCA GCC GAA GTG GCC GTA TAC CAG GAC GTG TAT GCT GTA CAT GCA 563 Ala Ala Glu Val Ala Val Tyr Gln Asp Val Tyr Ala Val His Ala 145 150 155												
CCA ACA TCG CTG TAC CAT CAG GCG ATG AAA GGT GTC AGA ACG GCG 608 Pro Thr Ser Leu Tyr His Gln Ala Met Lys Gly Val Arg Thr Ala 160 165 170												
TAT TGG ATT GGG TTT GAC ACC CCG TTT ATG TTT GAC GCG CTA 653 Tyr Trp Ile Gly Phe Asp Thr Thr Pro Phe Met Phe Asp Ala Leu 175 180 185												

GATGGCGGAT GTGTGACATA CACGACGCCA AAAGATTTTG TTCCAGCTCC TGCCACCTCC 60

Figure 5 (2)

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GCG Ala																		1058
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GAA Glu																		1103
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TGT Cys																		1148
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GAC																		1193
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Figure 5 (3)

AAG	GC2	A GA	CCT	r gar	r gat	C GA	AAA A	CC	CIX	G GG	r GT	CG2	GAC	AGG	1328
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														Met	
415					420		•			425			•		
														GTG	1418
H15		Me	с Туз	Lys	Lys 435		Asp	Thr	GIL	440		· Val	Lys	Val	
430					433	'				340	•				
CCT	TCA	GA	3 TTI	AAC	TCG	TTC	GTC	ATC	CCG	AGC	CTA	TGG	TCT	ACA	1463
Pro	Ser	Gli	ı Phe	Asn	Ser	Phe	Val	Ile	Pro	Ser	Leu	Trp	Ser	Thr	
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	• •			Pro											1300
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														TCA	1553
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GCC	AGG	GAT	CCI	GAA	CAA	GAG	GAG	AAG	GAG	AGG	TTG	GAG	GCC	GAG	1598
	Arg	Asp	Ala	Glu		Glu	Glu	Lys	Glu	_	Leu	Glu	Ala	Glu	
490					495					500					
CTG	ACT	AGA	GAA	GCC	TTA	CCA	ccc	CTC	GTC	ccc	ATC	GCG	CCG	GCG.	1643
				Ala											
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030	300	<i>a</i> a3	OTTO	OMC.	CAC	cenc .	CAC	com	C2.2	C2.2	CM3	030	mam.	G) G	1688
				Val											1000
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														GTC	1733 ⁻
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333					540					515					
ACC	GCA	CAG	CCG	AAC	GAC	GTA	CTA	CTA	GGA	AAT	TAC	GTA	GTT	CTG	1778
Thr .	Ala	Gln	Pro	Asn		Val	Leu	Leu	G1y		Tyr	Val	Val	Leu	
550					555					560					
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Ser															
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Figure 5 (4)

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			Val Pr			TTG AGC GA	
						AAC AGG AA Asn Arg Lys	
						ACC GAC GAC Thr Asp Glu	
						GCC GAG TAC	
						GAG GAA GCG Glu Glu Ala	
Ser Gly I 685	eu Val	Leu Val 690	Gly Glu	Leu Thi	r Asn Pro 695	CCG TTC CAT Pro Phe His	
						GCA CCA TAT Ala Pro Tyr	2228
Lys Thr T 715	hr Val	Val Gly 720	Val Phe	Gly Val	Pro Gly 725	TCA GGC AAG Ser Gly Lys	
Ser Ala I 730	le Ile	Lys Ser 735	Leu Val	Thr Lys	His Asp 740	CTG GTC ACC Leu Val Thr	
Ser Gly Ly 745	ys Lys (Glu Asn 750	Cys Gln	Glu Ile	Val Asn 755	GAC GTG AAG Asp Val Lys	2363
Lys His A 760	rg Gly 1	Lys Gly 1 765	Thr Ser	Arg Glu	Asn Ser . 770	GAC TCC ATC Asp Ser Ile	2408
						TAT GTG GAC Tyr Val Asp	2453
						CTA ATT GCT Leu Ile Ala	2498
						SAC CCC AAG	2543

Figure 5 (5)

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CAA TGC GGA TTC TTC AAT ATG ATG CAG CTT AAG GTG AAC TTC AAC 2588 Gln Cys Gly Phe Phe Asn Met Met Gln Leu Lys Val Asn Phe Asn 820 825 830	}
CAC AAC ATC TGC ACT GAA GTA TGT CAT AAA AGT ATA TCC AGA CGT 2633 His Asn Ile Cys Thr Glu Val Cys His Lys Ser Ile Ser Arg Arg 835 840 845	
TGC ACG CGT CCA GTC ACG GCC ATC GTG TCT ACG TTG CAC TAC GGA 2678 Cys Thr Arg Pro Val Thr Ala Ile Val Ser Thr Leu His Tyr Gly 850 855 860	
GGC AAG ATG CGC ACG ACC AAC CCG TGC AAC AAA CCC ATA ATC ATA 2723 Gly Lys Met Arg Thr Thr Asn Pro Cys Asn Lys Pro Ile Ile Ile 865 870 875	
GAC ACC ACA GGA CAG ACC AAG CCC AAG CCA GGA GAC ATC GTG TTA 2768 Asp Thr Thr Gly Gln Thr Lys Pro Lys Pro Gly Asp Ile Val Leu 880 885 890	
ACA TGC TTC CGA GGC TGG GCA AAG CAG CTG CAG TTG GAC TAC CGT 2813 Thr Cys Phe Arg Gly Trp Ala Lys Gln Leu Gln Leu Asp Tyr Arg 895 900 905	
GGA CAC GAA GTC ATG ACA GCA GCA GCA TCT CAG GGC CTC ACC CGC 2858 Gly His Glu Val Met Thr Ala Ala Ala Ser Gln Gly Leu Thr Arg 910 915 920	
AAA GGG GTA TAC GCC GTA AGG CAG AAG GTG AAT GAA AAT CCC TTG 2903 Lys Gly Val Tyr Ala Val Arg Gln Lys Val Asn Glu Asn Pro Leu 925 930 935	
TAT GCC CCT GCG TCG GAG CAC GTG AAT GTA CTG CTG ACG CGC ACT 2948 Tyr Ala Pro Ala Ser Glu His Val Asn Val Leu Leu Thr Arg Thr 940 945 950	
GAG GAT AGG CTG GTG TGG AAA ACG CTG GCC GGC GAT CCC TGG ATT 2993 Glu Asp Arg Leu Val Trp Lys Thr Leu Ala Gly Asp Pro Trp Ile 955 960 965	
AAG GTC CTA TCA AAC ATT CCA CAG GGT AAC TTT ACG GCC ACA TTG 3038 Lys Val Leu Ser Asn Ile Pro Gln Gly Asn Phe Thr Ala Thr Leu 970 975 980	
GAA GAA TGG CAA GAA GAA CAC GAC AAA ATA ATG AAG GTG ATT GAA 3083 Glu Glu Trp Gln Glu Glu His Asp Lys Ile Met Lys Val Ile Glu 985 990 995	
GGA CCG GCT GCG CCT GTG GAC GCG TTC CAG AAC AAA GCG AAC GTG 3128 Gly Pro Ala Ala Pro Val Asp Ala Phe Gln Asn Lys Ala Asn Val 1000 1005 1010	
TGT TGG GCG AAA AGC CTG GTG CCT GTC CTG GAC ACT GCC GGA ATC 3173 Cys Trp Ala Lys Ser Leu Val Pro Val Leu Asp Thr Ala Gly Ile 1015 1020 1025	

Figure 5 (6)

10/33

Arg Leu				ı Glı	ı Tr				e Il	e Th				
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Thr Lys 1060	Ty	г Туі	r Gly		_) Let	ı Asp	Sei			ı Phe	: Se	c Ala	
CCG AAG	GTY	TC(CTG	TAT	TAC	GAG	AAC	AAC	CAC	TGG	GAT	· AAC	: AGA	3353
Pro Lys 1075	Va:	l Sei	: Leu	_	_	Glu	a Asn	Asr		-) Asp) Asr	Arg	
CCT GGT	GG?	A AGO	ATG	TAI	GGA	TTC	: AAT	GCC	GCA	ACA	GCI	GCC	AGG	3398
Pro Gly 1090	Gly	/ Arc			_	Phe	Asn	Ala			Ala	Ala	Arg	
CTG GAA	GCI	AGA	CAT	ACC	TTC	CIG	AAG	GGG	CAG	TGG	CAT	ACG	GGC	3443
Leu Glu 1105	Ala	Arg			Phe	Leu	Lys	Gly		_	His	Thr	Gly	
														3488
Lys Gln 1120	Ala	Val			Glu	Arg	Lys				Leu	Ser	Val	٠
														3533
Leu Asp 1135	Asn	Val			Ile	Asn	Arg			Pro	His	Ala	Leu	
GTG GCT	GAG	TAC	AAG	ACG	GTT	AAA	GGC	AGT	AGG	GTT	GAG	TGG	CTG	3578
Val Ala 1150	Glu	Tyr	_		Val	Lys	Gly		-	Val	Glu	Trp	Leu	
GTC AAT	AAA	GTA	aga	GGG	TAC	CAC	GTC	CTG	CTG	GTG	AGT	GAG	TAC	3623
Val Asn 1165	Lys	Val	-		Tyr	His	Val			Val	Ser	Glu	Tyr	
AAC CTG	GCT	TTG	CCT	CGA	CGC	AGG	GTC	ACT	TGG	TTG	TCA	CCG	CTG	3668
Asn Leu 1180	Ala	Leu			Arg	Arg	Va1			Leu	Ser	Pro	Leu	
AAT GTC	ACA	GGC	GCC	GAT	AGG	TGC	TAC	GAC	CTA	agt	TTA	GGA	CTG	3713
	Thr	Gly		_	Arg	Суѕ	Tyr	_		Ser	Leu	Gly	Leu	
1133			1	200					1203					
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1210	usb	WTG	_		rne	vab	nen			val .	AS []	TTG	uis	
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Thr Glu I 1225	Phe	Arg			His	Tyr	Gln (_	Val .	Asp 1	His .	Ala	
	Arg Leu 1030 GAG GAC GAU Asp 1045 ACC AAG Thr Lys 1060 CCG AAG Pro Lys 1075 CCT GGT Pro Gly 1090 CTG GAA Leu Glu 1105 AAG CAG Lys Gln 1120 CTG GAC Leu Asp 1135 GTG GCT Val Ala 1150 GTC AAT Val Asn 1165 AAC CTG ASN Leu 1180 AAT GTC ASN Val 1195 CCG GCT GAT CASN Val Asn 1195 CCG GCT GAT CASN Val ASN CTG ASN Val ASN CTG GCT GCT ASN CTG GCT ASN CTG GCT GCT ASN CTG GCT GCT ASN CTG GCT GCT GCT ASN CTG GCT GCT GCT GCT ASN CTG GCT GCT GCT GCT GCT GCT GCT GCT GCT	Arg Leu Th 1030 GAG GAC AG Glu Asp Ar 1045 ACC AAG TA Thr Lys Ty 1060 CCG AAG GTA Pro Lys Va 1075 CCT GGT GGA Pro Gly Gly 1090 CTG GAA GCT Leu Glu Ala 1105 AAG CAG GCA Lys Gln Ala 1120 CTG GAC AAT Leu Asp Asn 1135 GTG GCT GAG Val Ala Glu 1150 GTC AAT AAA Val Asn Lys 1165 AAC CTG GCT ASN Leu Ala 1180 AAT GTC ACA Asn Val Thr 1195 CCG GCT GAC Pro Ala Asp 1210 ACG GAA TTC Thr Glu Phe	Arg Leu Thr Aldings GAG GAC AGA GCC Glu Asp Arg Aldings 1045 ACC AAG TAC TACT Thr Lys Tyr Tyr 1060 CCG AAG GTG TCC Pro Lys Val Ser 1075 CCT GGT GGA AGC Pro Gly Gly Arg 1090 CTG GAA GCT AGA Leu Glu Ala Arg 1105 AAG CAG GCA GTT Lys Gln Ala Val 1120 CTG GAC AAT GTA Leu Asp Asn Val 1135 GTG GCT GAG TAC Val Ala Glu Tyr 1150 GTC AAT AAA GTA Val Asn Lys Val 1165 AAC CTG GCT TTG Asn Leu Ala Leu 1180 AAT GTC ACA GGC Asn Val Thr Gly 1195 CCG GCT GAC GCC Pro Ala Asp Ala 1210 ACG GAA TTC AGA Thr Glu Phe Arg	Arg Leu Thr Ala Glu 1030 GAG GAC AGA GCT TAG Glu Asp Arg Ala Tyr 1045 ACC AAG TAC TAT GGA Thr Lys Tyr Tyr Gly 1060 CCG AAG GTG TCC CTG Pro Lys Val Ser Leu 1075 CCT GGT GGA AGG ATG Pro Gly Gly Arg Met 1090 CTG GAA GCT AGA CAT Leu Glu Ala Arg His 1105 AAG CAG GCA GTT ATC Lys Gln Ala Val Ile 1120 CTG GAC AAT GTA ATT Leu Asp Asn Val Ile 1135 GTG GCT GAG TAC AAG Val Ala Glu Tyr Lys 1150 GTC AAT AAA GTA AGA Val Asn Lys Val Arg 1165 AAC CTG GCT TTG CCT Asn Leu Ala Leu Pro 1180 AAT GTC ACA GGC GCC Asn Val Thr Gly Ala 1195 CCG GCT GAC GCC GGC Pro Ala Asp Ala Gly 1210 ACG GAA TTC AGA ATC Thr Glu Phe Arg Ile	Arg Leu Thr Ala Glu Glu 1030 GAG GAC AGA GCT TAC TCC Glu Asp Arg Ala Tyr Sei 1045 ACC AAG TAC TAT GGA GTT Thr Lys Tyr Tyr Gly Val 1060 CCG AAG GTG TCC CTG TAT Pro Lys Val Ser Leu Tyr 1075 CCT GGT GGA AGG ATG TAT Pro Gly Gly Arg Met Tyr 1090 CTG GAA GCT AGA CAT ACC Leu Glu Ala Arg His Thr 1105 AAG CAG GCA GTT ATC GCA Lys Gln Ala Val Ile Ala 1120 CTG GAC AAT GTA ATT CCT Leu Asp Asn Val Ile Pro 1135 CTG GCT GAG TAC AAG ACG Val Ala Glu Tyr Lys Thr 1150 AAC CTG GCT TTG CCT CGA Asn Leu Ala Leu Pro Arg 1180 AAT GTC ACA GGC GCC GAT Asn Val Thr Gly Ala Asp 1195 CCG GAT GAC GCC GCC AGG Pro Ala Asp Ala Gly Arg 1210 CCCG GCT GAC GCC GCC AGG Pro Ala Asp Ala Gly Arg 1210 CCCG GAA TTC AGA ATC CAC Thr Glu Phe Arg Ile His	Arg Leu Thr Ala Glu Glu Tr 1030 1035 GAG GAC AGA GCT TAC TCT CCC Glu Asp Arg Ala Tyr Ser Pro 1045 1050 ACC AAG TAC TAT GGA GTT GAC Thr Lys Tyr Tyr Gly Val Asp 1060 1065 CCG AAG GTG TCC CTG TAT TAC Pro Lys Val Ser Leu Tyr Tyr 1075 1080 CCT GGT GGA AGG ATG TAT GGA Pro Gly Gly Arg Met Tyr Gly 1090 1095 CTG GAA GCT AGA CAT ACC TTC Leu Glu Ala Arg His Thr Phe 1105 1110 AAG CAG GCA GTT ATC GCA GAA Lys Gln Ala Val Ile Ala Glu 1120 1125 CTG GAC AAT GTA ATT CCT ATC Leu Asp Asn Val Ile Pro Ile 1135 1140 GTG GCT GAG TAC AAG ACG GTT Val Ala Glu Tyr Lys Thr Val 1150 1155 GTC AAT AAA GTA AGA GGG TAC Val Asn Lys Val Arg Gly Tyr 1165 1170 AAC CTG GCT TTG CCT CGA CGC Asn Leu Ala Leu Pro Arg Arg 1180 1185 AAT GTC ACA GGC GCC GAT AGG Asn Val Thr Gly Ala Asp Arg 1195 1200 CCG GCT GAC GCC GGC AGG TTC Pro Ala Asp Ala Gly Arg Phe 1210 1215 ACG GAA TTC AGA ATC CAC CAC Thr Glu Phe Arg Ile His His	Arg Leu Thr Ala Glu Glu Trp Set 1030 1035 GAG GAC AGA GCT TAC TCT CCA GTG Glu Asp Arg Ala Tyr Ser Pro Vai 1045 1050 ACC AAG TAC TAT GGA GTT GAC CTG Thr Lys Tyr Tyr Gly Val Asp Let 1060 1065 CCG AAG GTG TCC CTG TAT TAC GAC Pro Lys Val Ser Leu Tyr Tyr Gly Phe 1075 1080 CCT GGT GGA AGG ATG TAT GGA TTC Pro Gly Gly Arg Met Tyr Gly Phe 1090 1095 CTG GAA GCT AGA CAT ACC TTC CTG Leu Glu Ala Arg His Thr Phe Leu 1105 1110 AAG CAG GCA GTT ATC GCA GAA AGA Lys Gln Ala Val Ile Ala Glu Arg 1120 1125 CTG GAC AAT GTA ATT CCT ATC AAC Leu Asp Asn Val Ile Pro Ile Asn 1135 1140 GTG GCT GAG TAC AAG ACG GTT AAA Val Ala Glu Tyr Lys Thr Val Lys 1150 1155 GTC AAT AAA GTA AGA GGG TAC CAC Val Asn Lys Val Arg Gly Tyr His 1165 1170 AAC CTG GCT TTG CCT CGA CGC AGG Asn Leu Ala Leu Pro Arg Arg Arg 1180 1185 AAT GTC ACA GGC GCC GAT AGG TGC Asn Val Thr Gly Ala Asp Arg Cys 1195 1200 CCG GCT GAC GCC GGC AGG TTC GAC Pro Ala Asp Ala Gly Arg Phe Asp 1210 1215 ACG GAA TTC AGA ATC CAC CAC TAC Thr Glu Phe Arg Ile His His Tyr	Arg Leu Thr Ala Glu Glu Trp Ser The 1030 1035 GAG GAC AGA GCT TAC TCT CCA GTG GTG GTG GIU Asp Arg Ala Tyr Ser Pro Val Val 1045 1050 ACC AAG TAC TAT GGA GTT GAC CTG GAC Thr Lys Tyr Tyr Gly Val Asp Leu Asp 1060 1065 CCG AAG GTG TCC CTG TAT TAC GAG AAC Pro Lys Val Ser Leu Tyr Tyr Glu Asn 1075 1080 CCT GGT GGA AGG ATG TAT GGA TTC AAT Pro Gly Gly Arg Met Tyr Gly Phe Asn 1090 1095 CTG GAA GCT AGA CAT ACC TTC CTG AAG Leu Glu Ala Arg His Thr Phe Leu Lys 1105 1110 AAG CAG GCA GTT ATC GCA GAA AGA AAA Lys Gln Ala Val Ile Ala Glu Arg Lys 1120 1125 CTG GAC AAT GTA ATT CCT ATC AAC CGC Leu Asp Asn Val Ile Pro Ile Asn Arg 1135 1140 GTG GCT GAG TAC AAG ACG GTT AAA GGC Val Ala Glu Tyr Lys Thr Val Lys Gly 1150 1155 GTC AAT AAA GTA AGA GGG TAC CAC GTC Val Asn Lys Val Arg Gly Tyr His Val 1165 1170 AAC CTG GCT TTG CCT CGA CGC AGG GTC Asn Leu Ala Leu Pro Arg Arg Arg Val 1180 1185 AAT GTC ACA GGC GCC GAT AGG TCC TAC Asn Val Thr Gly Ala Asp Arg Cys Tyr 1195 1200 CCG GCT GAC GCC GGC AGG TTC GAC TTG CTG GAA ATC AAC CAC GTC CAC GTC ASn Val Thr Gly Ala Asp Arg Cys Tyr 1195 1200 CCG GCT GAC GCC GCC AGG TTC GAC TTG CTG GAA TTC AGA ATC CAC CAC TAC CAG GTC GAA ATC CAC CAC TAC CAG GTC CAC GAA ATC CAC CAC TAC CAG GTC GAA ATC CAC CAC TAC CAG GTC CAC GAA TTC AGA ATC CAC CAC TAC CAG GTC CAC GAA TTC AGA ATC CAC CAC TAC CAG GTC CAC GAA TTC AGA ATC CAC CAC TAC CAG GTC CAC GAA TTC AGA ATC CAC CAC TAC CAC GTC TTC GIU Phe Arg Ile His His Tyr Gln GTC TAC THE GIU Phe Arg Ile His His Tyr Gln GTC TAC TAC GAA TTC CAC CAC TAC CAC TAC CAC GTC TTC GIU Phe Arg Ile His His Tyr Gln GTC TAC TAC GAA TTC CAC CAC TAC CAC GTC TTC GIU Phe Arg Ile His His Tyr Gln GTC TAC TAC GAA TTC CAC CAC TAC CAC TAC CAC TAC CAC TAC CAC GTC TTC GIU Phe Arg Ile His His Tyr Gln GTC TAC TAC TAC CAC GTC TTC GAC TTG CAC TTC GAC	Arg Leu Thr Ala Glu Glu Trp Ser Thr II 1030 1035 GAG GAC AGA GCT TAC TCT CCA GTG GTG GCG Glu Asp Arg Ala Tyr Ser Pro Val Val Ala 1045 1050 ACC AAG TAC TAT GGA GTT GAC CTG GAC AG Thr Lys Tyr Tyr Gly Val Asp Leu Asp Ser 1060 1065 CCG AAG GTG TCC CTG TAT TAC GAG AAC AAC Pro Lys Val Ser Leu Tyr Tyr Glu Asn Asr 1075 1080 CCT GGT GGA AGG ATG TAT GGA TTC AAT GCC Pro Gly Gly Arg Met Tyr Gly Phe Asn Ala 1090 1095 CTG GAA GCT AGA CAT ACC TTC CTG AAG GCG Leu Glu Ala Arg His Thr Phe Leu Lys Gly 1110 AAG CAG GCA GTT ATC GCA GAA AGA AAA ATC Lys Gln Ala Val IIe Ala Glu Arg Lys IIe 1120 1125 CTG GAC AAT GTA ATT CCT ATC AAC CGC AGG Leu Asp Asn Val IIe Pro IIe Asn Arg Arg 1135 1140 GTG GCT GAG TAC AAG ACG GTT AAA GGC AGT Val Ala Glu Tyr Lys Thr Val Lys Gly Ser 1150 1155 GTC AAT AAA GTA AGA GGG TAC CAC GTC CTG Val Asn Lys Val Arg Gly Tyr His Val Leu 1165 1170 AAC CTG GCT TTG CCT CGA CGC AGG GTC ACT Asn Leu Ala Leu Pro Arg Arg Arg Val Thr 1180 1185 AAT GTC ACA GGC GCC GAT AGG TGC TAC GAC Asn Val Thr Gly Ala Asp Arg Cys Tyr Asp 1195 1200 CCG GCT GAC GCC GCC AGG TTC GAC TTG GTC Pro Ala Asp Ala Gly Arg Phe Asp Leu Val 1210 1215 ACG GAA TTC AGA ATC CAC CAC TAC CAG CAG Thr Glu Phe Arg IIe His His Tyr Gln Gln	Arg Leu Thr Ala Glu Glu Trp Ser Thr Ile II 1030 1035 104 GAG GAC AGA GCT TAC TCT CCA GTG GTG GCC TR Glu Asp Arg Ala Tyr Ser Pro Val Val Ala Let 1045 1050 1055 ACC AAG TAC TAT GGA GTT GAC CTG GAC AGT GGC Thr Lys Tyr Tyr Gly Val Asp Leu Asp Ser Gly 1060 1065 1076 CCG AAG GTG TCC CTG TAT TAC GAG AAC AAC CAC Pro Lys Val Ser Leu Tyr Tyr Glu Asn Asn His 1075 1080 1085 CCT GGT GGA AGG ATG TAT GGA TTC AAT GCC GCA Pro Gly Gly Arg Met Tyr Gly Phe Asn Ala Ala 1090 1095 1100 CTG GAA GCT AGA CAT ACC TTC CTG AAG GGG CAG Leu Glu Ala Arg His Thr Phe Leu Lys Gly Glu 11105 1110 AAG CAG GCA GTT ATC GCA GAA AGA AAA ATC CAA Lys Gln Ala Val Ile Ala Glu Arg Lys Ile Glu 1120 1125 1130 CTG GAC AAT GTA ATT CCT ATC AAC CGC AGG CTG Leu Asp Asn Val Ile Pro Ile Asn Arg Arg Leu 1135 1140 1145 GTG GCT GAG TAC AAG ACG GTT AAA GGC AGT AGG Val Ala Glu Tyr Lys Thr Val Lys Gly Ser Arg 1150 1155 1160 GTC AAT AAA GTA AGA GGG TAC CAC GTC CTG CTG Val Asn Lys Val Arg Gly Tyr His Val Leu Leu 1165 1170 1175 AAC CTG GCT TTG CCT CGA CGC AGG GTC ACT TGG Asn Leu Ala Leu Pro Arg Arg Arg Val Thr Trp 1180 1185 1190 AAT GTC ACA GGC GCC GAT AGG TTC GAC TTC GTA Asn Val Thr Gly Ala Asp Arg Cys Tyr Asp Leu 1195 1200 1205 CCG GCT GAC GCC GGC AGG TTC GAC CTG CTG TTP CO Ala Asp Ala Gly Arg Phe Asp Leu Val Phe 1210 1225 ACG GAA TTC AGA ATC CAC CAC TAC CAC CTG TTT Tro Ala Asp Ala Gly Arg Phe Asp Leu Val Phe 1210 1225 ACG GAA TTC AGA ATC CAC CAC TAC CAC CAC TTT Thr Glu Phe Arg Ile His His Tyr Gln Gln Cys	Arg Leu Thr Ala Glu Glu Trp Ser Thr Ile Ile The 1030 1035 1040 GAG GAC AGA GCT TAC TCT CCA GTG GTG GCC TTG AAG GIU Asp Arg Ala Tyr Ser Pro Val Val Ala Leu Ash 1045 1050 1055 ACC AAG TAC TAT GGA GTT GAC CTG GAC AGT GGC CTG Thr Lys Tyr Tyr Gly Val Asp Leu Asp Ser Gly Leu 1060 1065 1070 CCG AAG GTG TCC CTG TAT TAC GAG AAC AAC CAC TGG Pro Lys Val Ser Leu Tyr Tyr Glu Ash Ash His Trp 1075 1080 1085 CCT GGT GGA AGG ATG TAT GGA TTC AAT GCC GCA ACA Pro Gly Gly Arg Met Tyr Gly Phe Ash Ala Ala Thr 1090 1095 1100 CTG GAA GCT AGA CAT ACC TTC CTG AAG GGG CAG TGG Leu Glu Ala Arg His Thr Phe Leu Lys Gly Gln Trp 1105 1110 1115 AAG CAG GCA GTT ATC GCA GAA AGA AAA ATC CAA CCG Lys Gln Ala Val Ile Ala Glu Arg Lys Ile Gln Pro 1120 1125 1130 CTG GAC AAT GTA ATT CCT ATC AAC CGC AGG CTG CCG Leu Asp Ash Val Ile Pro Ile Ash Arg Arg Leu Pro 1135 1140 1145 GTG GCT GAG TAC AAG ACG GTT AAA GGC AGT AGG GTT Val Ala Glu Tyr Lys Thr Val Lys Gly Ser Arg Val 1150 1155 1160 GTC AAT AAA GTA AGA GGG TAC CAC CAC CTC CTG CTG GTG CAL AAA GTA AAA GTA AAA GTA GGG TTC AAT AAA GGC AAT GTA AGA GGG TAC CAC CTC CTG CTG GTG CAL AAT AAA GTA AAA GTA AAA GGC AAT GTA ATT CCT ATC AAC CAC GTC CTG CTG GTG CAL AAT AAA GTA AAA GTA AAA GGC AAT AGA AGA CAG GTT AAA GGC AAT AGG GTT AAT AAA GTA AAA GTA AAA GGC AAT AGA GGG TAC CAC CTC CTG CTG GTG CAL AAT AAA GTA AAA GTA AGA GGG TAC CAC CTC CTG CTG GTG CAL AAT AAA GTA AAA GAG AGG GTC ACT TGG TTG AAT AAA GTA AAA GTA AAA AGA GGG TAC CAC CTC CTG CTG GTG AAT AAA GTA AAA GAA AAA AATC CAC CAC CTC CTG CTG GTG AAT AAA ATC CAC GCC GAT AGG GTC ACT TGG TTG AAT AAA GTA AAA AATA AAA GAA AAA A	Arg Leu Thr Ala Glu Glu Trp Ser Thr Ile Ile Thr Ala 1030 1035 1040 GAG GAC AGA GCT TAC TCT CCA GTG GTG GCC TTG AAT GAZ GLU ASP ARG Ala Tyr Ser Pro Val Val Ala Leu Asn Glu 1045 1050 1055 ACC AAG TAC TAT GGA GTT GAC CTG GAC AGT GGC CTG TTT Thr Lys Tyr Tyr Gly Val Asp Leu Asp Ser Gly Leu Phe 1060 1065 1070 CCG AAG GTG TCC CTG TAT TAC GAG AAC AAC CAC TCG GAT Pro Lys Val Ser Leu Tyr Tyr Glu Asn Asn His Trp Asp 1075 1080 1085 CCT GGT GGA AGG ATG TAT GGA TTC AAT GCC GCA ACA GCT Pro Gly Gly Arg Met Tyr Gly Phe Asn Ala Ala Thr Ala 11090 1095 1100 CTG GAA GCT AGA CAT ACC TTC CTG AAG GGG CAG TGG CAT Leu Glu Ala Arg His Thr Phe Leu Lys Gly Gln Trp His 1105 1110 1115 AAG CAG GCA GTT ATC GCA GAA AGA AAA ATC CAA CCG CTT Lys Gln Ala Val Ile Ala Glu Arg Lys Ile Gln Pro Leu 1120 1125 1130 CTG GAC AAT GTA ATT CCT ATC AAC GCC AGG CTG CCG CAC Leu Asp Asn Val Ile Pro Ile Asn Arg Arg Leu Pro His 1135 1140 CTG GAC TAC AAG ACG GTT AAT CCT ATC AAC CCC AGG CTG CCG CAC Leu Asp Asn Val Ile Pro Ile Asn Arg Arg Leu Pro His 1135 1140 GTG GCT GAG TAC AAG ACG GTT AAA GCC AGT AGG GTT GAG Val Ala Glu Tyr Lys Thr Val Lys Gly Ser Arg Val Glu 1155 1160 GTC AAT AAA GTA AGA AGG GTA CAC CTC CTG CTG GTG AGT Val Asn Lys Val Arg Gly Tyr His Val Leu Leu Val Ser 1165 1170 1175 AAC CTG GCT TTG CCT CGA CGC AGG GTC ACT TGG TTG TCA Asn Leu Ala Leu Pro Arg Arg Arg Val Thr Trp Leu Ser 1185 1190 AAT GTC ACA GCC GCC GAT AGG TCC TAC GAC CTA AGT TTA Asn Val Thr Gly Ala Asp Arg Cys Tyr Asp Leu Ser Leu 1195 1200 1205 CCC GCT GAC GCC GCC AGG TTC GAC TTG GTC TTT GTG AAC Pro Ala Asp Ala Gly Arg Phe Asp Leu Val Phe Val Asp 1210 1215	Arg Leu Thr Ala Glu Glu Trp Ser Thr Ile Ile Thr Ala Ph 1030 1035 1040 GAG GAC AGA GCT TAC TCT CCA GTG GTG GCC TTG AAT GAA AT GIU Asp Arg Ala Tyr Ser Pro Val Val Ala Leu Asn Glu Il. 1045 1050 1055 ACC AAG TAC TAT GGG GTT GAC CTG GAC AGT GGC CTG TTT TCT. Thr Lys Tyr Tyr Gly Val Asp Leu Asp Ser Gly Leu Phe Ser 1060 1065 1070 CCG AAG GTG TCC CTG TAT TAC GAG AAC AAC CAC TGG GAT AAC Pro Lys Val Ser Leu Tyr Tyr Glu Asn Asn His Trp Asp Asr 1075 1080 1085 CCT GGT GGA AGG ATG TAT GGA TTC AAT GCC GCA ACA GCT GCC Pro Gly Gly Arg Met Tyr Gly Phe Asn Ala Ala Thr Ala Ala 1090 1095 1100 CTG GAA GCT AGA CAT ACC TTC CTG AAG GGG CAG TGG CAT ACC Leu Glu Ala Arg His Thr Phe Leu Lys Gly Gln Trp His Thr 1110 1115 AAG CAG GCA GTT ATC GCA GAA AGA AAA ATC CAA CCG CTT TCT Lys Gln Ala Val Ile Ala Glu Arg Lys Ile Gln Pro Leu Ser 1120 1125 CTG GAC AAT GTA ATT CCT ATC AAC CGC AGG CTG CCG CAC CCC Leu Asp Asn Val Ile Pro Ile Asn Arg Arg Leu Pro His Ala 1135 1140 1145 CTG GCT GAG TAC AAG ACG GTT AAA GGC AGT AGG GTT GAG TGG CTG CTG GAT GAG TAC GAT AGA GCT GCT GCT GAT GAG TGC GCT GCT GCT GCT GAT GAG TGC GCT GCT GCT GCT GCT GAT GAG TGC GCT GCT GCT GCT GCT GCT GCT GCT GC	GAG GAC AGA GCT TAC TCT CCA GTG GTG GCC TTG AAT GAA ATT TGC Glu Asp Arg Ala Tyr Ser Pro Val Val Ala Leu Asn Glu Ile Cys 1045 1050 1055 ACC AAG TAC TAT GGA GTT GAC CTG GAC AGT GGC CTG TTT TCT GCC Thr Lys Tyr Tyr Gly Val Asp Leu Asp Ser Gly Leu Phe Ser Ala 1060 1065 1070 CCG AAG GTG TCC CTG TAT TAC GAG AAC AAC CAC TGG GAT AAC AGA Pro Lys Val Ser Leu Tyr Tyr Glu Asn Asn His Trp Asp Asn Arg 1075 1080 CCT GGT GGA AGG ATG TAT GGA TTC AAT GCC GCA ACA GCT GCC AGG Pro Gly Gly Arg Met Tyr Gly Phe Asn Ala Ala Thr Ala Ala Arg 1090 1095 1100 CTG GAA GCT AGA CAT ACC TTC CTG AAG GGG CAG TGG CAT ACG GGC Leu Glu Ala Arg His Thr Phe Leu Lys Gly Gln Trp His Thr Gly 1105 1110 AAG CAG GCA GTT ATC GCA GAA AGA AAA ATC CAA CCG CTT TCT GTG Lys Gln Ala Val Ile Ala Glu Arg Lys Ile Gln Pro Leu Ser Val 1120 CTG GAC AAT GTA ATT CCT ATC AAC CGC AGG CTG CCG CAC GCC CTG Leu Asp Asn Val Ile Pro Ile Asn Arg Arg Leu Pro His Ala Leu 1135 1140 GTG GCT GAG TAC AAG ACG GTT AAA GGC AGT AGG GTT GAG TGG CTG Val Ala Glu Tyr Lys Thr Val Lys Gly Ser Arg Val Glu Trp Leu 1155 1160 GTC AAT AAA GTA AAG ACG GTT AAA GGC AGT AGG GTT GAG TGG CTG Val Ala Glu Tyr Lys Thr Val Lys Gly Ser Arg Val Glu Trp Leu 1155 1160 GTC AAT AAA GTA AGA GGG TAC CAC GTC CTC CTG GTG AGT GAG TGG CTG GAC TTG CCT CGA CGC AGG GTC ACT TGG TTG TGA GTG ASn Leu Ala Leu Pro Arg Arg Arg Val Thr Trp Leu Ser Pro Leu 1185 1170 AAC CTG GCT TTG CCT CGA CGC AGG GTC ACT TGG TTG TCA CCG CTG Asn Leu Ala Leu Pro Arg Arg Arg Val Thr Trp Leu Ser Pro Leu 1180 1185 1190 AAT GTC ACA GGC GCC GAT AGG TTC TAC GAC CTA AGT TTA GGA CTG Asn Val Thr Gly Ala Asp Arg Cys Tyr Asp Leu Ser Leu Gly Leu 1195 1200 1205 ACG GCT GAG TAC AGA ATC CAC CAC TTG CTG GTC TTG GAC CAC GCC CTG CAC GCC GAC AGG TTC GAC TTG GTC TTG GAC AAT CAC Pro Ala Asp Ala Gly Arg Phe Asp Leu Val Phe Val Asn Ile His 1210 12215 1220 ACG GAA TTC AGA ATC CAC CAC TAC CAC CAG TTG GTC GAC CAC GCC Thr Glu Phe Arg Ile His His Tyr Gln Gln Cys Val Asp His Ala

Figure 5 (7)

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ATG ANG CTG CAG ATG CTT GGG GGA GAT GCG CTA CC	SA CIG CIA AAA 3848
Met Lys Leu Gln Met Leu Gly Gly Asp Ala Leu An	rg Leu Leu Lys
1240 1245 1250	
CCC GGC GGC ATC TTG ATG AGA GCT TAC GGA TAC GC	CC GAT AAA ATC 3893
Pro Gly Gly Ile Leu Met Arg Ala Tyr Gly Tyr Al	la Asp Lys Ile
1255 1260 1265	•
AGC GAA GCC GTT GTT TCC TCC TTA AGC AGA AAG TT	
Ser Glu Ala Val Val Ser Ser Leu Ser Arg Lys Ph	e Ser Ser Ala
1270 1275 1280	
AGA GTG TTG CGC CCG GAT TGT GTC ACC AGC AAT AC	A GAA GTG TTC 3983
Arg Val Leu Arg Pro Asp Cys Val Thr Ser Asn Th	r Glu Val Phe
1285 1290 1295	
TTG CTG TTC TCC AAC TTT GAC AAC GGA AAG AGA CC	
Leu Leu Phe Ser Asn Phe Asp Asn Gly Lys Arg Pr	o Ser Thr Leu
1300 1305 1310	
CAC CAG ATG AAT ACC AAG CTG AGT GCC GTG TAT GC	C GGA GAA GCC 4073
His Gln Met Asn Thr Lys Leu Ser Ala Val Tyr Ala	a Gly Glu Ala
1315 1320 1325	
ATG CAC ACG GCC GGG TGT GCA CCA TCC TAC AGA GT	T AAG AGA GCA 4118
Met His Thr Ala Gly Cys Ala Pro Ser Tyr Arg Val	l Lys Arg Ala
1330 1335 1340	
GAC ATA GCC ACG TGC ACA GAA GCG GCT GTG GTT AAC	GCA GCT AAC 4163
Asp Ile Ala Thr Cys Thr Glu Ala Ala Val Val Asr	n Ala Ala Asn
1345 1350 1355	
GCC CGT GGA ACT GTA GGG GAT GGC GTA TGC AGG GCC	
Ala Arg Gly Thr Val Gly Asp Gly Val Cys Arg Ala	Val Ala Lys
1360 1365 1370	
AAA TGG CCG TCA GCC TTT AAG GGA GCA GCA ACA CCA	GTG GGC ACA 4253
Lys Trp Pro Ser Ala Phe Lys Gly Ala Ala Thr Pro	Val Gly Thr
1375 1380 1385	
ATT AAA ACA GTC ATG TGC GGC TCG TAC CCC GTC ATC	CAC GCT GTA 4298
Ile Lys Thr Val Met Cys Gly Ser Tyr Pro Val Ile	His Ala Val
1390 1395 1400	
ecg cct aat tic ict gcc acg act gaa gcg gaa ggg	
Ala Pro Asn Phe Ser Ala Thr Thr Glu Ala Glu Gly	Asp Arg Glu
1405 1410 1415	
TG GCC GCT GTC TAC CGG GCA GTG GCC GCC GAA GTA	
eu Ala Ala Val Tyr Arg Ala Val Ala Ala Glu Val	Asn Arg Leu
1425 1430	
CA CTG AGC AGC GTA GCC ATC CCG CTG CTG TCC ACA	GGA GTG TTC 4433
er Leu Ser Ser Val Ala Ile Pro Leu Leu Ser Thr	Gly Val Phe
435 1440 1445	
SUBSTITUTE SHEET	• •.

ATG AAG CTG CAG ATG CTT GGG GGA GAT GCG CTA CGA CTG CTA AAA 3848

Figure 5 (8)

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AGC GGC GGA AGA GAT AGG CTG CAG CAA TCC CTC AAC CAT CTA TTC 4478 Ser Gly Gly Arg Asp Arg Leu Gln Gln Ser Leu Asn His Leu Phe 1450 1455 1460
ACA GCA ATG GAC GCC ACG GAC GCT GAC GTG ACC ATC TAC TGC AGA 4523 Thr Ala Met Asp Ala Thr Asp Ala Asp Val Thr Ile Tyr Cys Arg 1465 1470 1475
GAC AAA AGT TGG GAG AAG AAA ATC CAG GAA GCC ATT GAC ATG AGG 4568 Asp Lys Ser Trp Glu Lys Lys Ile Gln Glu Ala Ile Asp Met Arg 1480 1485 1490
ACG GCT GTG GAG TTG CTC AAT GAT GAC GTG GAG CTG ACC ACA GAC 4613 Thr Ala Val Glu Leu Leu Asn Asp Asp Val Glu Leu Thr Thr Asp 1495 1500 1505
TTG GTG AGA GTG CAC CCG GAC AGC AGC CTG GTG GGT CGT AAG GGC 4658 Leu Val Arg Val His Pro Asp Ser Ser Leu Val Gly Arg Lys Gly 1510 1515 1520
TAC AGT ACC ACT GAC GGG TCG CTG TAC TCG TAC TTT GAA GGT ACG 4703 Tyr Ser Thr Thr Asp Gly Ser Leu Tyr Ser Tyr Phe Glu Gly Thr 1525 1530 1535
AAA TTC AAC CAG GCT GCT ATT GAT ATG GCA GAG ATA CTG ACG TTG 4748 Lys Phe Asn Gln Ala Ala Ile Asp Met Ala Glu Ile Leu Thr Leu 1540 1545 1550
TGG CCC AGA CTG CAA GAG GCA AAC GAA CAG ATA TGC CTA TAC GCG 4793 Trp Pro Arg Leu Gln Glu Ala Asn Glu Gln Ile Cys Leu Tyr Ala 1555 1560 1565
CTG GGC GAA ACA ATG GAC AAC ATC AGA TCC AAA TGT CCG GTG AAC 4838 Leu Gly Glu Thr Met Asp Asn Ile Arg Ser Lys Cys Pro Val Asn 1570 1757 1580
GAT TCC GAT TCA TCA ACA CCT CCC AGG ACA GTG CCC TGC CTG TGC 4883 Asp Ser Asp Ser Ser Thr Pro Pro Arg Thr Val Pro Cys Leu Cys 1585 1590 1595
CGC TAC GCA ATG ACA GCA GAA CGG ATC GCC CGC CTT AGG TCA CAC 4928 Arg Tyr Ala Met Thr Ala Glu Arg Ile Ala Arg Leu Arg Ser His 1600 1605 1610
CAA GTT AAA AGC ATG GTG GTT TGC TCA TCT TTT CCC CTC CCG AAA 4973 Gln Val Lys Ser Met Val Val Cys Ser Ser Phe Pro Leu Pro Lys 1615 1620 1625
TAC CAT GTA GAT GGG GTG CAG AAG GTA AAG TGC GAG AAG GTT CTC 5018 Tyr His Val Asp Gly Val Gln Lys Val Lys Cys Glu Lys Val Leu 1630 1635 1640
CTG TTC GAC CCG ACG GTA CCT TCA GTG GTT AGT CCG CGG AAG TAT 5063 Leu Phe Asp Pro Thr Val Pro Ser Val Val Ser Pro Arg Lys Tyr 1645 1650 1655

Figure 5 (9)

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														G TTI y Phe	
166					1665		, mo	a mar	n mo	167				n 100	. E1E3
	Leu					: Asp					r Ala			r ACC Thr	
	Ser					Glr					as e			TAC Tyr	5198
	Pro					Val					Val			GAA Glu	5243
	Ala					Leu					. His			CCC Pro	5288
	Asp			Ąsp				CCG Pro			Pro	-		CCG Pro	5333
	Arg			Tyr				CGC Arg			Glu			GTG Val	5378
	Ala			Lys				GCC Ala	Pro		Thr			AGG Arg	5423
	Lys			Leu			_	GAC Asp	Phe		Glu				5468
	Ala			Ser				TTC Phe	Gly					GTC Val	5513
	Arg			Arg				TAT Tyr	Ile						5558
	Ser			Leu				TCC Ser	Va1						5603
CAG (Gln (1840	Cys	GCA Ala	CAA Gln	Leu	GAT Asp .845	GCG Ala	GTC Val	CAG Gln	Glu	GAG Glu L850	AAA Lys	ATG Met	TAC Tyr	CCG Pro	5648
				Thr	Glu 860	Arg	Glu	AAG Lys	Leu 1	Leu 1865					5693
					S	UBS	TIT	JTE	SHE	E I					

Figure 5 (10)

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CAG ATO Gln Met 1870		Ser				Tyr		
AAA GTO Lys Val 1885		Met				Arg		
GGG GCC Gly Ala 1900		Tyr				Arg		5828
TAC GCG Tyr Ala 1915		Tyr			Tyr	Pro		5873
GAA AGA Glu Arg 1930		Ser			Ile			5918
TAC CTA Tyr Leu 1945		Asn '			Ala			5963
GAT GAA Asp Glu 1960		Ala '			Val			6008
TGC TTG Cys Leu 1975		Ala 1			Ala			6053
CCG AAA Pro Lys 1990		la 1			Thr			6098
CCG TCA Pro Ser 2005		ln A			Asn		 	 6143
ACC AAG Thr Lys 2020			sn V		Met .			6188
ATG GAC Met Asp 2035			he A		ys			6233
GC TCC (Cys Ser (2050			rp G		la 1			6278
ATA ACC I le Thr 1 2065			le T		al 1			6323

Figure 5 (11)

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Pı					u Ph				s As		TT CC	
Le					t As				l As		AA CG 's Arg	
GA As	T GI			тсс	A GG o Gl			C AC	A GA r Gl		A CCC	
Ly					n Ala				ı Al		T TAC a Tyr	
	u Cy				g Glu				J Let		T GTG a Val	
	u Ar				Thr			ATG	TCC		A GAC 1 Asp	6593
	e Asp				Ser				Gly		GTT Val	6638
СТА	A GAG) Ile	GCA	TCA Ser		Lys	AGC	CAG Gln		TCC Ser	6683
TTG	GCT Ala		GGI Gly	TTA	ATG		GAA Glu	GAT	CTA		GAT Asp	6728
CAG	TAC		GAC	TTG			GCC Ala	TTT				6773
AGC	TGT Cys		CCA Pro	ACT			TTC Phe	aag				6818
ATG	AAA Lys		ATG Met	TTT			TTT Phe	ATT				6863
AAC	ATC Ile		GCA Ala	AGC			GAG Glu	CAG				6908
TCC	GCC Ala		GCC Ala	TTC			GAC . Asp .	AAC .				6953

Figure 5 (12)

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GTG ATC TCC GAC AAG CTG ATG GCG GAG AGG TGC GCG TCG TGG GTC 6 Val Ile Ser Asp Lys Leu Met Ala Glu Arg Cys Ala Ser Trp Val 2290 2295 2300	998
AAC ATG GAG GTG AAG ATC ATT GAC GCT GTC ATG GGC GAA AAA CCC 70 Asn Met Glu Val Lys Ile Ile Asp Ala Val Met Gly Glu Lys Pro 2305 2310 2315	043
CCA TAT TTT TGT GGG GGA TTC ATA GTT TTT GAC AGC GTC ACA CAG 70 Pro Tyr Phe Cys Gly Gly Phe Ile Val Phe Asp Ser Val Thr Gln 2320 2325 2330)88
100 000 mgg ggm ggm mg1 010 001 ggm 110 001	.33
GGT AAG CCG CTA ACA GCT GAA GAC AAG CAG GAC GAA GAC AGG CGA 71 Gly Lys Pro Leu Thr Ala Glu Asp Lys Gln Asp Glu Asp Arg Arg 2350 2355 2360	78
CGA GCA CTG AGT GAC GAG GTT AGC AAG TGG TTC CGG ACA GGC TTG 72. Arg Ala Leu Ser Asp Glu Val Ser Lys Trp Phe Arg Thr Gly Leu 2365 2370 2375	23
GGG GCC GAA CTG GAG GTG GCA CTA ACA TCT AGG TAT GAG GTA GAG 720 Gly Ala Glu Leu Glu Val Ala Leu Thr Ser Arg Tyr Glu Val Glu 2380 2385 2390	58
GGC TGC AAA AGT ATC CTC ATA GCC ATG ACC ACC TTG GCG AGG GAC 731 Gly Cys Lys Ser Ile Leu Ile Ala Met Thr Thr Leu Ala Arg Asp 2395 2400 2405	13
ATT AAG GCG TTT AAG AAA TTG AGA GGA CCT GTT ATA CAC CTC TAC 735 Ile Lys Ala Phe Lys Lys Leu Arg Gly Pro Val Ile His Leu Tyr 2410 2415 2420	8
GGC GGT CCT AGA TTG GTG CGT TAA TACACAGAAT TCTGATTATA GCGCACTATT Gly Gly Pro Arg Leu Val Arg 2425 2430	7412
ATAGCACC ATG AAT TAC ATC CCT ACG CAA ACG TTT TAC GGC CGC CGG 74 Met Asn Tyr Ile Pro Thr Gln Thr Phe Tyr Gly Arg Arg 5 10	59
TGG CGC CCG CGC CCG GCC GCC CGT CCT TGC CCG TTG CAG GCC ACT 750. Trp Arg Pro Arg Pro Ala Ala Arg Pro Trp Pro Leu Gln Ala Thr 15 20 25	4 .
CCG GTG GCT CCC GTC CCC GAC TTC CAG GCC CAG CAG ATG CAG 7549 Pro Val Ala Pro Val Val Pro Asp Phe Gln Ala Gln Gln Met Gln 30 35 40)
CAA CTC ATC AGC GCC GTA AAT GCG CTG ACA ATG AGA CAG AAC GCA 7594 Gln Leu Ile Ser Ala Val Asn Ala Leu Thr Met Arg Gln Asn Ala	ŀ

Figure 5 (13)

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				Pro P				AG AAG A Ys Lys Ly 70		CC 7639
				Thr G				C AAC GO le Asn Gl 85		CG 7684 hr
Gln G				Lys As				AC AAG AA PP Lys Ly 100		AG 7729 Ys
Lys P					g Met			G ATT GA s Ile Gl 115		
	le Ph				s Glu			C ACT GG 1 Thr Gly 130		
TGC CT Cys Le 13	u Va	G GGC 1 Gly	GAC A	AAA GT Lys Va 14	1 Met	Lys Lys	CCT GC Pro Ala	C CAC GTO a His Val 145	G AAA GG l Lys Gl	A 7864 Y
	e Ası				ı Ala			TTC AAC A Phe Lys 160		
	з Туг				a Ala			Val His		
	p Ala				His			GAG GGA Glu Gly 190		
	o His				Gln			GGT AGG Gly Arg 205		
ATA CCC Ile Pro 210	Thr	GGA Gly	GCG GG Ala G	GC AAA ly Lys 215	Pro	GGA G	AC AGT Ser	GGC CGG Gly Arg 220	CCC ATC	8089
TTT GAC Phe Asp 225	Asn	AAG Lys	GGG AG	GG GTA g Val 230	GTC (GCT A Ala I	TC GTC le Val	CTG GGC Leu Gly 235	GGG GCC Gly Ala	8134
AAC GAG Asn Glu 240	GGC Gly	TCA (CGC AC Arg Th	A GCA ir Ala 245	CTG :	ICG G Ser V	TG GTC al Val	ACC TGG Thr Trp 250	AAC AAA Asn Lys	8179
	Val							GAA GAG Glu Glu 265		8224

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Figure 5 (14)

GCC	: ccc	CI	G AI	T AC	T GC	C ATY	G TG	T GT	CCT	T GC	C AA	r GC	r ac	C TTC	8269
Ala	Pro	Le	u Il	e Th	r Ala	a Met	t Cy	s Vai	l Le	u Ala	a Ası	n Ala	a Th	r Phe	1
	270	1			•	279	5				280)			
CCG	TGC	TT	C CA	G CC	c ccc	G TGT	r GT	A CC	r TG	C TG	C TAT	r gaz	AA	C AAC	8314
Pro	Cys	Ph	e Gl	n Pr	o Pro	Cys	Va.	l Pro	Cy:	s Cys	з Туг	: Glu	ı Ası	n Asn	
	285					290)				295	5			
GCA	GAG	GC	C AC	A CT	A CGC	ATC	CIX	GAC	GA?	r aac	GTG	GAT	AGO	G CCA	8359
Ala	Glu	Al	a Th	r Le	u Arg	Met	: Let	ı Glu	ı Ası) Asr	ı Val	Asp	Arg	g Pro	
	300					305	5				310) _			
GGG	TAC	TA	C GA	CT	CTI	CAG	GC2	A GCC	TTO	ACC	TGC	CGA	AAC	GGA	8404
														Gly	
	315	-				320					325	_			
ACA	AGA	CA	CG	G CG(AGC	GTG	TCG	CAA	CAC	TTC	: AAC	GTG	TAT	' AAG	8449
Thr	Arg	Hi	s Ar	g Arg	, Ser	. Val	Ser	Gln	His	Phe	Asn	Val	Tyr	Lys	
	330					335					340		_	_	
GCT	ACA	CGG	cc'	TAC	ATC	GCG	TAC	TGC	GCC	GAC	TGC	GGA	GCA	GGG	8494
Ala	Thr	Arg	j Pro	э Туг	: Ile	Ala	Tyr	Cys	Ala	Asp	Cys	Gly	Ala	Gly	
	345			_		350	_	_			355	_			
CAC	TCG	TG?	CA!	r ago	: ccc	GTA	GCA	ATT	GAA	GCG	GTC	AGG	TCC	GAA	8539
His	Ser	Cys	Hi	s Ser	Pro	Val	Ala	Ile	Glu	Ala	Val	Arg	Ser	Glu	
	360	-				365					370	Ī			
GCT	ACC	GAC	GGC	3 ATG	CTG	AAG	ATT	CAG	TTC	TCG	GCA	CAA	ATT	GCC	8584
Ala	Thr	Asp	Gly	Met	Leu	Lys	Ile	Gln	Phe	Ser	Ala	Gln	Ile	Gly	
	375	_				380					385			_	
ATA	GAT	AAG	AG1	GAC	AAT	CAT	GAC	TAC	ACG	AAG	ATA	AGG	TAC	GCA	8629
Ile	Asp	Lys	Ser	: Asp	Asn	His	Asp	Tyr	Thr	Lys	Ile	Arg	Tyr	Ala	
	390					395					400				
GAC	GGG	CAC	GCC	ATT	GAG	AAT	GCC	GTC	CGG	TCA	TCT	TTG	AAG	GTA	8674
Asp	Gly	His	Ala	Ile	Glu	Asn	Ala	Va1	Arg	Ser	Ser	Leu	Lys	Val	
	405					410					415				
					TGT										8719
Ala	Thr	Ser	Gly	Asp	Суз	Phe	Val	His	Gly	Thr	Met	Gly	His	Phe	
	420					425					430				
ATA	CTG	GCA	AAG	TGC	CCA	CCG	GGT	GAA	TTC	CTG	CAG	GTC	TCG	ATC	8764
Ile	Leu	Ala	Lys	Cys	Pro	Pro	Gly	Glu	Phe	Leu	Gln	Val	Ser	Ile	
	435					440					445				
CAG	GAC	ACC	AGA	AAC	GCG	GTC	CGT	GCC	TGC	AGA	ATA	CAA	TAT	CAT	8809
Gln	Asp	Thr	Arg	Asn	Ala	Val	Arg	Ala	Cys	Arg	Ile	Gln	Tyr	His	•
	4 50		-			455	-				460				
CAT	GAC	CCT	CAA	CCG	GTG	GGT	AGA	GAA	AAA	TTT	ACA	ATT	AGA	CCA	8854
His	Asp	Pro	Gln	Pro	Val	Gly	Arg	Glu	Lys	Phe	Thr	Ile	Arg	Pro	
	465					470					475				

Figure 5 (15)

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														C ACA		
His	480		ly Ly	ys Gl	u Il	e Pro 48:		s Th	r Th	r Ty:	r Gl 49		n Th	r Thr		
														T ACG	8944	
ALC	495		L V.	11 G1	u (31)	500		y Mei	L ni	s ne	50		O AS	p m		
														G ATC s Ile	8989	
	510					515					520					
														A ACC y Thr	9034	
	525					530)				535	5				
														TGT Cys	9079	
02,	540			J		545		nop	1100		550			. cys		
														TGG	9124	
Leu	555		n GI	п суз	s nis	560		vai	inr	Asp	565	-	. Lys	Trp		
														AGA	9169	
GIN	570	ASI	1 Se	r Pro) Pne	575	PIO	Arg	Ala	Asp	580		A A L	Arg		
				C CAT l His										TGC	9214	
5 10	585		, vu			590	20	110	Deu	nop	595	116	****	Cys		
				G GCG : Ala											9259	
3	600				3	605					610	3		9		
				CAC His										TCC Ser	9304	
	615					620		-			625					
				GGT Gly										GTG	9349	
	630	1111	Deu	GLY	GIU	635	FIO	GIII	171	urs	640	GIU	пр	Val		
				GAA Glu										ATG Met	9394	
	645				9	650			·		655	-~P	1			
				GGA Gly											9439	
	660			1		665					670	P				
				GGG Gly											9484	
	675		JIG	1	_	680		~ - 3			685	~=**		- ~4		
										A 15						

Figure 5 (16)

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CAC	TAC	C TA	C TA	T GG	G CT	TA(c ccc	GC	C GC	T AC	A GT	A TC	C GC	G GTC	9529
Glr		_	T Ty	r Gl	y Le	_		Ala	a Al	a Th			r Al	a Val	
	690)				69	5				70	0			
GTC	. යය	דג ב	YS AG	с тт	а сту	GCC	3 TTC	: ATZ	TC	G AT	מידי כ	c GCC	: TC	G TGC	9574
		-												r Cys	
	705	•				710					719			•	
														r gct	
Тут			u Va	l Ala	a Ale			Lys	Cys	s Let			Ty	r Ala	
	720	,				725	,				730	,			
TTA	ACA	CC	A GG	A GC	r GCA	GTI	r ccc	TGG	ACC	CIX	GGG	ATA	CTO	TGC	9664
Leu	Thr	Pr	o G1	y Ala	a Ala	. Val	Pro	Trp	Thi	: Let	ı Gly	Ile	Lev	ı Cys	
	735					740)				745	;			
maa	000		c co					3 CM	- CMC				3000		9709
														Ala	9/09
C _J S	750		O AL	,		755		561	741	. Alt	760		Met	, AIG	
															9754
Tyr			b yai	Glr	Asn			Leu	Phe	Trp			Phe	Ala	
	765					770					775				
GCC	CCT	GT.	r gcc	TGC	ATC	CTC	ATC	ATC	ACG	TAT	TGC	CTC	AGA	AAC	9799
				Cys											
	780					785					790				
СПС	CITC	m~n	n mee	יייביעיי	אמי	NCC.	COURT	an~an	- CALALIA	on on the same	CIT/C	C/D3	~	AGC	9844
				Cys											7044
	795		•			800					805				
														CCG	9889
Leu	810	ALE	Thr	Ala	Arg	815	ıyr	GIU	Hls	Ser	1717 820	Vai	Met	Pro	
	010					010					020				
AAC	GTG	GTG	GGG	TTC	CCG	TAT	AAG	GCT	CAC	ATT	GAA	AGG	CCA	GGA	9934
Asn		Val	Gly	Phe	Pro	_	Lys	Ala	His	Ile		Arg	Pro	Gly	
	825					830					835		•		
ጥልጥ	AGC	ccc	ריזיר	ACT	عكامة	CAG	DAKS.	CAG	بلملت	Cum	CAA	ACC) AGC	CTC	9979
				Thr											3373
_	840					845					850				
														_	
				AAT											10024
	855	ınr	Leu	Asn	Leu	860	туг	11e	Thr	Cys	865	TYE	гÀз	Inr	
	000														
GTC	GTC	CCG	TCG	CCG	TAC	GTG	AAG	TGC	TGC	GGC	GCC	TCA ·	GAG	TGC	10069
		Pro	Ser	Pro			Lys	Cys	Cys	Gly		Ser	Glu	Cys	
1	870					875					880				
ጥርር	ልርጥ	מממ	GNG	AAG	ርር ሞ	CAC	ጥልሮ ፣	CAA	ጥርጉ	ממ	، بادلت	ጥልሮ	מרא	CCC	10114
				Lys											70114
	885	, _				890				_	895			-4	
												_			

Figure 5 (17)

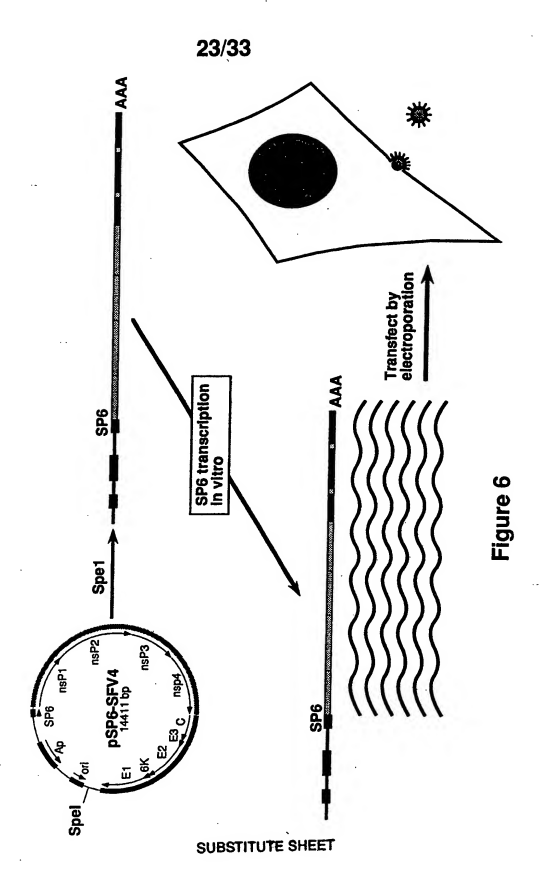
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	Pro			Gly			Cys	C TCA p Ser	
	Thi			Ala			g Sei	C GTA Val	
				Ala			Thr	A TCG A Ser	
							Asn	ACT Thr	10294
								GGT Gly	10339
Thr			GGG Gly			•		TTC Phe	10384
Asp			GTG Val	-	 	Val	 	 GAC	10429
Phe			Ser			Arg		ATC Ile	10474
Gln .			Glu			Tyr		GCA Ala	10519
Leu :			Pro			Val			10564
Thr (GGG Gly 1			Leu	 	 	10609
Thr A			AAG Lys 1			Cys (10654
Asn I			ATG Met 1			Gly i			10699
Ser M			GAC Asp :			Arg 1			10744

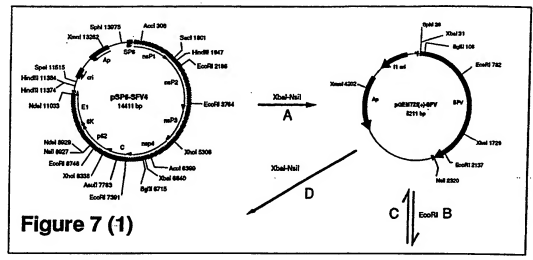
Figure 5 (18)

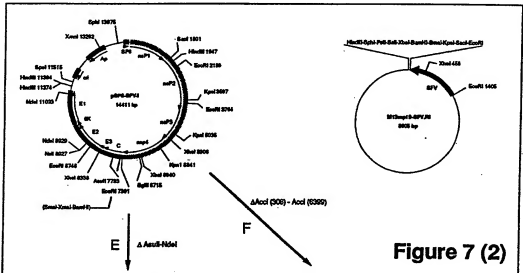
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CCG ACC ATC ATT GAC CTG ACT TGC ACA GTG GCT ACC TGT ACG CAC 10789 Pro Thr Ile Ile Asp Leu Thr Cys Thr Val Ala Thr Cys Thr His 1110 1115 1120
TCC TCG GAT TTC GGC GGC GTC TTG ACA CTG ACG TAC AAG ACC AAC 10834 Ser Ser Asp Phe Gly Gly Val Leu Thr Leu Thr Tyr Lys Thr Asn 1125 1130 1135
AAG AAC GGG GAC TGC TCT GTA CAC TCG CAC TCT AAC GTA GCT ACT 10879 Lys Asn Gly Asp Cys Ser Val His Ser His Ser Asn Val Ala Thr 1140 1145 1150
CTA CAG GAG GCC ACA GCA AAA GTG AAG ACA GCA GGT AAG GTG ACC 10924 Leu Gln Glu Ala Thr Ala Lys Val Lys Thr Ala Gly Lys Val Thr 1155 1160 1165
TTA CAC TTC TCC ACG GCA AGC GCA TCA CCT TCT TTT GTG GTG TCG 10969 Leu His Phe Ser Thr Ala Ser Ala Ser Pro Ser Phe Val Val Ser 1170 1175 1180
CTA TGC AGT GCT AGG GCC ACC TGT TCA GCG TCG TGT GAG CCC CCG 11014 Leu Cys Ser Ala Arg Ala Thr Cys Ser Ala Ser Cys Glu Pro Pro 1185 1190 1195
AAA GAC CAC ATA GTC CCA TAT GCG GCT AGC CAC AGT AAC GTA GTG 11059 Lys Asp His Ile Val Pro Tyr Ala Ala Ser His Ser Asn Val Val 1200 1205 1210
TTT CCA GAC ATG TCG GGC ACC GCA CTA TCA TGG GTG CAG AAA ATC 11104 Phe Pro Asp Met Ser Gly Thr Ala Leu Ser Trp Val Gln Lys Ile 1215 1220 1225
TCG GGT GGT CTG GGG GCC TTC GCA ATC GGC GCT ATC CTG GTG CTG 11149 Ser Gly Gly Leu Gly Ala Phe Ala Ile Gly Ala Ile Leu Val Leu 1230 1235 1240
GTT GTG GTC ACT TGC ATT GGG CTC CGC AGA TAA GTTAGGGTAG 11192 Val Val Val Thr Cys Ile Gly Leu Arg Arg 1245 1250
GCAATGGCAT TGATATAGCA AGAAAATTGA AAACAGAAAA AGTTAGGGTA AGCAATGGCA 11252
TATAACCATA ACTGTATAAC TTGTAACAAA GCGCAACAAG ACCTGCGCAA TTGGCCCCGT 11312
GGTCCGCCTC ACGGAAACTC GGGGCAACTC ATATTGACAC ATTAATTGGC AATAATTGGA 11372
AGCTTACATA AGCTTAATTC GACGAATAAT TGGATTTTTA TTTTATTTTG CAATTGGTTT 11432
ТТААТАТТТС САААААААА АААААААААА АААААААА
AAAAAAAAAA AAAAAAAAA ACTAG . 11517



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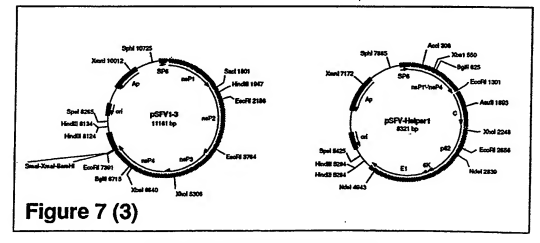
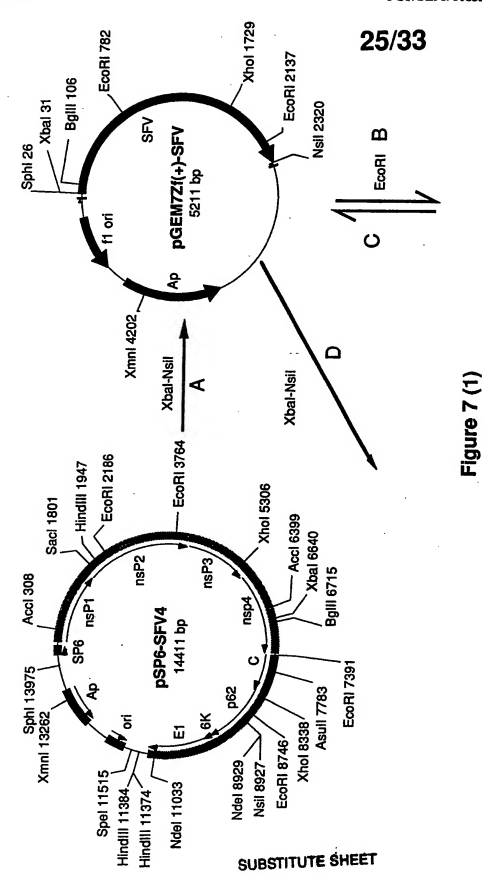
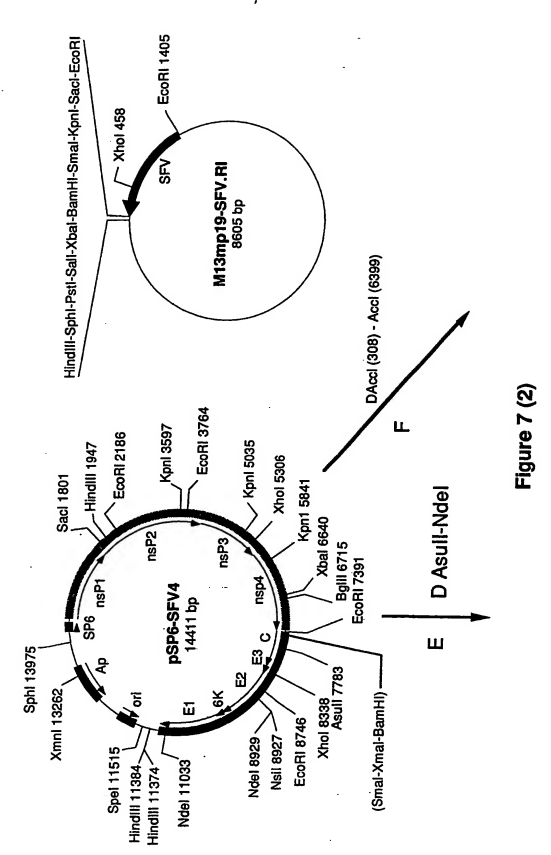


Figure 7 layout scheme





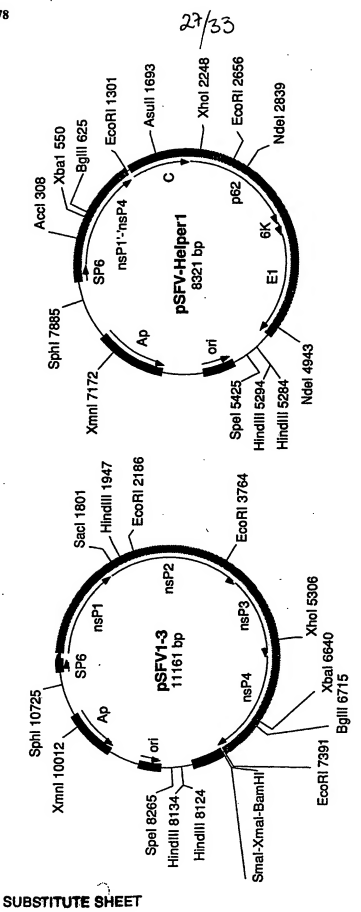


Figure 7 (3)

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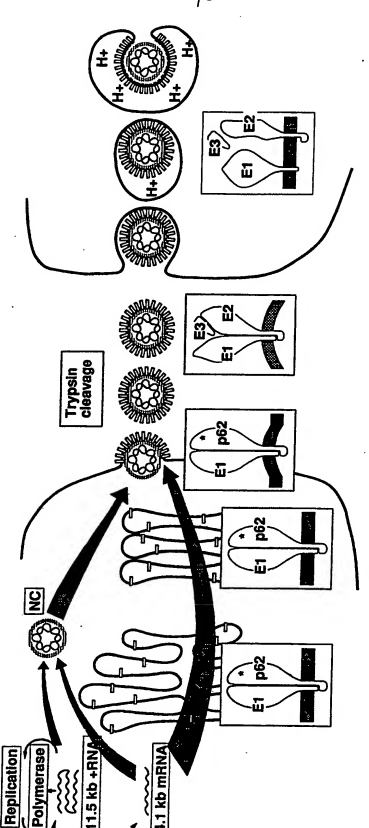


Figure 10

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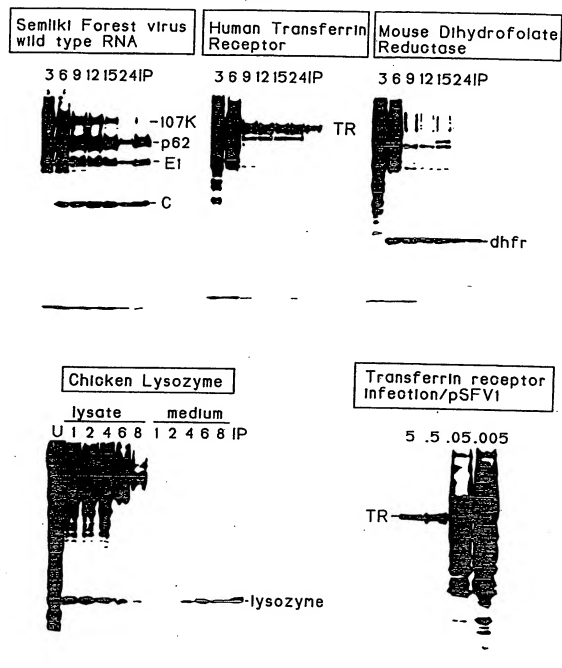


Figure 11

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	32/33		
GTC CAT SFV VECTOR E2 Val His SFV E2 SFV VECTOR E2	G HIV-gp120 HIV epitope		CCG GGC Pro SFV-HIV chimera
BamH1 GAT CC GAA CCG GCT AGA AAA GGC AAA GIU Pro Ala Arg Lys Gly Lys Asp 247	GGA AGA GCA TTT GTT GAG CCT TCT CGT AAA CAA CTC CTA Gly Arg Ala Phe Val Glu Asp	Cut with BamH1 Insert HIV oligo	GGA AGA GCA TTT GTT GAG GAT CCT TCT CGT AAA CAA CTC CTA Gly Arg Ala Phe Val Glu Asp
TCA CCT TTC GTC CCG AGA GCC GAC Ser Pro Phe Val Pro Arg Ala Asp Glu	GAT CCG CGT ATC CAG AGA GGA CCA GC GCA TAG GTC TCT CCT GGT ASP Pro Arg Ile Gln Arg Gly Pro		GAG GAT CCG CGT ATC CAG AGA GGA CCA CTC CTA GGC GCA TAG GTC TCT CCT GGT Glu Asp Pro Arg Ile Gln Arg Gly Pro
AAC Asn 			ga <u>f</u> crc Glu

Figure 12 (1)

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SFV spike

SFV virion

SUBSTITUTE SHEET

Figure 12 (2)

INTERNATIONAL SEARCH REPORT

International Application No PCT/SE 91/00855

-			International Application Ro PC1	/35 31/00000
I. CLAS	SIFICATIO	N OF SUBJECT MATTER (if several cia:	ssification symbols apply, indicate all) ⁸	
Accordin	g to Interna	tional Patent Classification (IPC) or to bot	h National Classification and IPC	
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			her than Minimum Documentation	
		to me extent that such Docume	ents are included in Fields Searched ⁸	
SE.DK.F	FI.NO c	lasses as above		
		NSIDERED TO BE RELEVANT®		
Category *	Citati	on of Document, ¹¹ with indication, where a	appropriate, of the relevant passages 12	Relevant to Claim No.13
X	PROC.N	ATL.ACAD.SCI., Vol. 84,	1987 (USA) Robin	1-2,5,9-
	Le	vis et al: "Engineered d	efective interfering	11,20-
	RN	As of Sindbis virus expr	ess bacterial	22,29,
	ch	loramphenicol acetyltran	sferase in avian	30
- 1	ce	lls", see page 4811 - p	age 4815	
- 1	es	pecially page 4811, colu	mn 1 lines	ł
ł	1-	4,12-15,46-48; column 2	lines 25-28 and	
	pa	ge 4812 column 1 lines 2	4-26 column 2	1
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Y [WO, A1	, 8912095 (APPLIED BIOTE	CHNOLOGY, INC.)	3,4,6-8,
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International Application No. PCT/SE 91/00855

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cit	Patent document ed in search report	Publication date	Paten men	t family nber(s)	Publication date		
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